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(54) Title: MICRORNA MOLECULES

(57) Abstract: In Caenorhabditis elegans, lin-4 and let-7 encode 22- and 21 -nucleotide RNAs, respectively, that function as key regulators of developmental timing. Because the appearance of these short RNAs is regulated during development, they are also referred to as "small temporal RNAs" (stRNAs). We show that many more 21- and 22-nt expressed RNAs, termed microRNAs, (miRNAs), exist in invertebrates and vertebrates, and that some of these novel RNAs, similar to let-7 stRNA, are also highly conserved. This suggests that sequence-specific post-transcriptional regulatory mechanisms mediated by small RNAs are more general than previously appreciated.





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## MicroRNA molecules

# Description

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The present invention relates to novel small expressed (micro)RNA molecules associated with physiological regulatory mechanisms, particularly in developmental control.

In Caenorhabditis elegans, lin-4 and let-7 encode 22- and 21-nucleotide RNAs, respectively (1, 2), that function as key regulators of developmental timing (3-5). Because the appearance of these short RNAs is regulated during development, they are also referred to as "microRNAs" (miRNAs) or small temporal RNAs (stRNAs) (6). lin-4 and let-21 are the only known miRNAs to date.

Two distinct pathways exist in animals and plants in which 21- to 23-nucleotide RNAs function as post-transcriptional regulators of gene expression. Small interfering RNAs (siRNAs) act as mediators of sequence-specific mRNA degradation in RNA interference (RNAi) (7-11) whereas miRNAs regulate developmental timing by mediating sequence-specific repression of mRNA translation (3-5). siRNAs and miRNAs are excised from double-stranded RNA (dsRNA) precursors by Dicer (12, 13, 29), a multidomain RNase III protein, thus producing RNA species of similar size. However, siRNAs are believed to be double-stranded (8, 11, 12), while miRNAs are single-stranded (6).

We show that many more short, particularly 21- and 22-nt expressed RNAs, termed microRNAs (miRNAs), exist in invertebrates and vertebrates, and that some of these novel RNAs, similar to let-7 RNA (6), are also highly conserved. This suggests that sequence-specific post-transcriptional

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regulatory mechanisms mediated by small RNAs are more general than previously appreciated.

The present invention relates to an isolated nucleic acid molecule comprising:

- (a) a nucleotide sequence as shown in Table 1, Table 2, Table 3 or Table 4
- (b) a nucleotide sequence which is the complement of (a),

99%, to a sequence of (a) or (b) and/or

(c) a nucleotide sequence which has an identity of at least 80%, preferably of at least 90% and more preferably of at least

(d) a nucleotide sequence which hybridizes under stringent conditions to a sequence of (a), (b) and/or (c).

In a preferred embodiment the invention relates to miRNA molecules and analogs thereof, to miRNA precursor molecules and to DNA molecules encoding miRNA or miRNA precursor molecules.

Preferably the identity of sequence (c) to a sequence of (a) or (b) is at least 90%, more preferably at least 95%. The determination of identity (percent) may be carried out as follows:

l = n : L

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wherein I is the identity in percent, n is the number of identical nucleotides between a given sequence and a comparative sequence as shown in Table 1, Table 2, Table 3 or Table 4 and L is the length of the comparative sequence. It should be noted that the nucleotides A, C, G and U as depicted in Tables 1, 2, 3 and 4 may denote ribonucleotides,

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deoxyribonucleotides and/or other nucleotide analogs, e.g. synthetic nonnaturally occurring nucleotide analogs. Further nucleobases may be substituted by corresponding nucleobases capable of forming analogous Hbonds to a complementary nucleic acid sequence, e.g. U may be substituted by T.

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Further, the invention encompasses nucleotide sequences which hybridize under stringent conditions with the nucleotide sequence as shown in Table 1, Table 2, Table 3 or Table 4, a complementary sequence thereof or a highly identical sequence. Stringent hybridization conditions comprise washing for 1 h in 1 x SSC and 0.1% SDS at 45°C, preferably at 48°C and more preferably at 50°C, particularly for 1 h in 0.2 x SSC and 0.1% SDS.

The isolated nucleic acid molecules of the invention preferably have a length of from 18 to 100 nucleotides, and more preferably from 18 to 80 nucleotides. It should be noted that mature miRNAs usually have a length of 19-24 nucleotides, particularly 21, 22 or 23 nucleotides. The miRNAs, however, may be also provided as a precursor which usually has a length of 50-90 nucleotides, particularly 60-80 nucleotides. It should be noted that the precursor may be produced by processing of a primary transcript which may have a length of >100 nucleotides.

The nucleic acid molecules may be present in single-stranded or double-stranded form. The miRNA as such is usually a single-stranded molecule, while the mi-precursor is usually an at least partially self-complementary molecule capable of forming double-stranded portions, e.g. stem- and loop-structures. DNA molecules encoding the miRNA and miRNA precursor molecules. The nucleic acids may be selected from RNA, DNA or nucleic acid analog molecules, such as sugar- or backbone-modified ribonucleotides or deoxyribonucleotides. It should be noted, however, that other nucleic analogs, such as peptide nucleic acids (PNA) or locked nucleic acids (LNA), are also suitable.

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In an embodiment of the invention the nucleic acid molecule is an RNA- or DNA molecule, which contains at least one modified nucleotide analog, i.e. a naturally occurring ribonucleotide or deoxyribonucleotide is substituted by a non-naturally occurring nucleotide. The modified nucleotide analog may be located for example at the 5'-end and/or the 3'-end of the nucleic acid molecule.

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Preferred nucleotide analogs are selected from sugar- or backbone-modified ribonucleotides. It should be noted, however, that also nucleobase-modified ribonucleotides, i.e. ribonucleotides, containing a non-naturally occurring nucleobase instead of a naturally occurring nucleobase such as uridines or cytidines modified at the 5-position, e.g. 5-(2-amino)propyl uridine, 5-bromo uridine; adenosines and guanosines modified at the 8-position, e.g. 8-bromo guanosine; deaza nucleotides, e.g. 7-deaza-adenosine; O- and N-alkylated nucleotides, e.g. N6-methyl adenosine are suitable. In preferred sugar-modified ribonucleotides the 2'-OH-group is replaced by a group selected from H, OR, R, halo, SH, SR, NH<sub>2</sub>, NHR, NR<sub>2</sub> or CN, wherein R is C<sub>1</sub>-C<sub>6</sub> alkyl, alkenyl or alkynyl and halo is F, Cl, Br or I. In preferred backbone-modified ribonucleotides the phosphoester group connecting to adjacent ribonucleotides is replaced by a modified group, e.g. of phosphothioate group. It should be noted that the above modifications may be combined.

The nucleic acid molecules of the invention may be obtained by chemical synthesis methods or by recombinant methods, e.g. by enzymatic transcription from synthetic DNA-templates or from DNA-plasmids isolated from recombinant organisms. Typically phage RNA-polymerases are used for transcription, such as T7, T3 or SP6 RNA-polymerases.

The invention also relates to a recombinant expression vector comprising a recombinant nucleic acid operatively linked to an expression control sequence, wherein expression, i.e. transcription and optionally further

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processing results in a miRNA-molecule or miRNA precursor molecule as described above. The vector is preferably a DNA-vector, e.g. a viral vector or a plasmid, particularly an expression vector suitable for nucleic acid expression in eukaryotic, more particularly mammalian cells. The recombinant nucleic acid contained in said vector may be a sequence which results in the transcription of the miRNA-molecule as such, a precursor or a primary transcript thereof, which may be further processed to give the miRNA-molecule.

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Further, the invention relates to diagnostic or therapeutic applications of the claimed nucleic acid molecules. For example, miRNAs may be detected in biological samples, e.g. in tissue sections, in order to determine and classify certain cell types or tissue types or miRNA-associated pathogenic disorders which are characterized by differential expression of miRNA-molecules or miRNA-molecule patterns. Further, the developmental stage of cells may be classified by determining temporarily expressed miRNA-molecules.

Further, the claimed nucleic acid molecules are suitable for therapeutic applications. For example, the nucleic acid molecules may be used as modulators or targets of developmental processes or disorders associated with developmental dysfunctions, such as cancer. For example, miR-15 and miR-16 probably function as tumor-suppressors and thus expression or delivery of these RNAs or analogs or precursors thereof to tumor cells may provide therapeutic efficacy, particularly against leukemias, such as B-cell chronic lymphocytic leukemia (B-CLL). Further, miR-10 is a possible regulator of the translation of Hox Genes, particularly Hox 3 and Hox 4 (or Scr and Dfd in Drosophila).

In general, the claimed nucleic acid molecules may be used as a modulator of the expression of genes which are at least partially complementary to said nucleic acid. Further, miRNA molecules may act as target for

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therapeutic screening procedures, e.g. inhibition or activation of miRNA molecules might modulate a cellular differentiation process, e.g. apoptosis.

Furthermore, existing miRNA molecules may be used as starting materials for the manufacture of sequence-modified miRNA molecules, in order to modify the target-specificity thereof, e.g. an oncogene, a multidrug-resistance gene or another therapeutic target gene. The novel engineered miRNA molecules preferably have an identity of at least 80% to the starting miRNA, e.g. as depicted in Tables 1, 2, 3 and 4. Further, miRNA molecules can be modified, in order that they are symetrically processed and then generated as double-stranded siRNAs which are again directed against therapeutically relevant targets.

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Furthermore, miRNA molecules may be used for tissue reprogramming procedures, e.g. a differentiated cell line might be transformed by expression of miRNA molecules into a different cell type or a stem cell.

For diagnostic or therapeutic applications, the claimed RNA molecules are preferably provided as a pharmaceutical composition. This pharmaceutical composition comprises as an active agent at least one nucleic acid molecule as described above and optionally a pharmaceutically acceptable carrier.

The administration of the pharmaceutical composition may be carried out by known methods, wherein a nucleic acid is introduced into a desired target cell in vitro or in vivo.

Commonly used gene transfer techniques include calcium phosphate, DEAE-dextran, electroporation and microinjection and viral methods [30, 31, 32, 33, 34]. A recent addition to this arsenal of techniques for the introduction of DNA into cells is the use of cationic liposomes [35].

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Commercially available cationic lipid formulations are e.g. Tfx 50 (Promega) or Lipofectamin 2000 (Life Technologies).

The composition may be in form of a solution, e.g. an injectable solution, a cream, ointment, tablet, suspension or the like. The composition may be administered in any suitable way, e.g. by injection, by oral, topical, nasal, rectal application etc. The carrier may be any suitable pharmaceutical carrier. Preferably, a carrier is used, which is capable of increasing the efficacy of the RNA molecules to enter the target-cells. Suitable examples of such carriers are liposomes, particularly cationic liposomes.

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Further, the invention relates to a method of identifying novel microRNA-molecules and precursors thereof, in eukaryotes, particularly in vertebrates and more particularly in mammals, such as humans or mice. This method comprises: ligating 5'- and 3'-adapter-molecules to the end of a size-fractionated RNA-population, reverse transcribing said adapter-ligated RNA-population, and characterizing said reverse transcribed RNA-molecules, e.g. by amplification, concatamerization, cloning and sequencing.

A method as described above already has been described in (8), however, for the identification of siRNA molecules. Surprisingly, it was found now that the method is also suitable for identifying the miRNA molecules or precursors thereof as claimed in the present application.

Further, it should be noted that as 3'-adaptor for derivatization of the 3'-OH group not only 4-hydroxymethylbenzyl but other types of derivatization groups, such as alkyl, alkyl amino, ethylene glycol or 3'-deoxy groups are suitable.

Further, the invention shall be explained in more detail by the following Figures and Examples:

#### Figure Legends

Fig. 1A. Expression of *D. melanogaster* miRNAs. Northern blots of total RNA isolated from staged populations of *D. melanogaster* were probed for the indicated miRNAs. The position of 76-nt val-tRNA is also indicated on the blots. 5S rRNA serves as loading control. E, embryo; L, larval stage; P, pupae; A, adult; S2, Schneider-2 cells. It should be pointed out, that S2 cells are polyclonal, derived from an unknown subset of embryonic tissues, and may have also lost some features of their tissue of origin while maintained in culture. miR-3 to miR-6 RNAs were not detectable in S2 cells (data not shown). miR-14 was not detected by Northern blotting and may be very weakly expressed, which is consistent with its cloning frequency. Similar miRNA sequences are difficult to distinguish by Northern blotting because of potential cross-hybridization of probes.

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Fig. 1B. Expression of vertebrate miRNAs. Northern blots of total RNA isolated from HeLa cells, mouse kidneys, adult zebrafish, frog ovaries, and S2 cells were probed for the indicated miRNAs. The position of 76-nt val-tRNA is also indicated on the blots. 5S rRNA from the preparations of total RNA from the indicated species is also shown. The gels used for probing of miR-18, miR-19a, miR-30, and miR-31 were not run as far as the other gels (see tRNA marker position). miR-32 and miR-33 were not detected by Northern blotting, which is consistent with their low cloning frequency. Oligodeoxynucleotides used as Northern probes were:

let-7a, 5 TACTATACAACCTACTACCTCAATTTGCC (SEQ ID NO:1);

let-7d, 5 'ACTATGCAACCTACTACCTCT (SEQ ID NO:2);

let-7e, 5 'ACTATACAACCTCCTACCTCA (SEQ ID NO:3);

D. melanogaster val-tRNA, 5 'TGGTGTTTCCGCCCGGGAA (SEQ ID NO:4);

miR-1, 5 'TGGAATGTAAAGAAGTATGGAG (SEQ ID NO:5);

miR-2b, 5 GCTCCTCAAAGCTGGCTGTGATA (SEQ ID NO:6);

miR-3, 5 'TGAGACACACTTTGCCCAGTGA (SEQ ID NO:7);

miR-4, 5 'TCAATGGTTGTCTAGCTTTAT (SEQ ID NO:8);

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miR-5, 5 CATATCACAACGATCGTTCCTTT (SEQ ID NO:9); miR-6, 5 ' AAAAAGAACAGCCACTGTGATA (SEQ ID NO:10); miR-7, 5 TGGAAGACTAGTGATTTTGTTGT (SEQ ID NO:11); miR-8, 5 'GACATCTTTACCTGACAGTATTA (SEQ ID NO:12); miR-9, 5 TCATACAGCTAGATAACCAAAGA (SEQ ID NO:13): 5 miR-10, 5 ACAAATTCGGATCTACAGGGT (SEQ ID NO:14); miR-11, 5 GCAAGAACTCAGACTGTGATG (SEQ ID NO:15); miR-12, 5 'ACCAGTACCTGATGTAATACTCA (SEQ ID NO:16); miR-13a, 5 'ACTCGTCAAAATGGCTGTGATA (SEQ ID NO:17); 10 miR-14, 5' TAGGAGAGAGAAAAGACTGA (SEQ ID NO:18): miR-15, 5 TAGCAGCACATAATGGTTTGT (SEQ ID NO:19); . . miR-16, 5 'GCCAATATTTACGTGCTGCTA (SEQ ID NO:20); miR-17, 5 TACAAGTGCCTTCACTGCAGTA (SEQ ID NO:21); miR-18, 5 TATCTGCACTAGATGCACCTTA (SEQ ID NO:22); miR-19a, 5 TCAGTTTTGCATAGATTTGCACA (SEQ ID NO:23); 15 miR-20, 5 TACCTGCACTATAAGCACTTTA (SEQ ID NO:24); miR-21, 5 'TCAACATCAGTCTGATAAGCTA (SEQ ID NO:25); miR-22, 5 'ACAGTTCTTCAACTGGCAGCTT (SEQ ID NO:26); miR-23, 5 'GGAAATCCCTGGCAATGTGAT (SEQ ID NO:27); miR-24, 5 CTGTTCCTGCTGAACTGAGCCA (SEQ ID NO:28); 20 miR-25, 5 TCAGACCGAGACAAGTGCAATG (SEQ ID NO:29); miR-26a, 5 'AGCCTATCCTGGATTACTTGAA (SEQ ID NO:30); miR-27; 5 AGCGGAACTTAGCCACTGTGAA (SEQ ID NO:31); miR-28, 5 CTCAATAGACTGTGAGCTCCTT (SEQ ID NO:32); miR-29, 5 'AACCGATTTCAGATGGTGCTAG (SEQ ID NO:33); 25 miR-30, 5 'GCTGCAAACATCCGACTGAAAG (SEQ ID NO:34); miR-31, 5 CAGCTATGCCAGCATCTTGCCT (SEQ ID NO:35); miR-32, 5' GCAACTTAGTAATGTGCAATA (SEQ ID NO:36); miR-33, 5' TGCAATGCAACTACAATGCACC (SEQ ID NO:37).

Fig. 2. Genomic organization of miRNA gene clusters. The precursor structure is indicated as box and the location of the miRNA within the

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precursor is shown in gray; the chromosomal location is also indicated to the right. (A) D. melanogaster miRNA gene clusters. (B) Human miRNA gene clusters. The cluster of let-7a-1 and let-7f-1 is separated by 26500 nt from a copy of let-7d on chromosome 9 and 17. A cluster of let-7a-3 and let-7b, separated by 938 nt on chromosome 22, is not illustrated.

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- Fig. 3. Predicted precursor structures of D. melanogaster miRNAs. RNA secondary structure prediction was performed using mfold version 3.1 [28] and manually refined to accommodate G/U wobble base pairs in the helical segments. The miRNA sequence is underlined. The actual size of the stemloop structure is not known experimentally and may be slightly shorter or longer than represented. Multicopy miRNAs and their corresponding precursor structures are also shown.
- Fig. 4. Predicted precursor structures of human miRNAs. For legend, see Fig. 3.
  - Fig. 5. Expression of novel mouse miRNAs. Northern blot analysis of novel mouse miRNAs. Total RNA from different mouse tissues was blotted and probed with a 5 '-radiolabeled oligodeoxynucleotide complementary to the indicated miRNA. Equal loading of total RNA on the gel was verified by ethidium bromide staining prior to transfer; the band representing tRNAs is shown. The fold-back precursors are indicated with capital L. Mouse brains were dissected into midbrain, mb, cortex, cx, cerebellum, cb. The rest of the brain, rb, was also used. Other tissues were heart, ht, lung, lg, liver, lv, colon, co, small intestine, si, pancreas, pc, spleen, sp, kidney, kd, skeletal muscle, sm, stomach, st, H, human Hela SS3 cells. Oligodeoxynucleotides used as Northern probes were:

miR-1a, CTCCATACTTCTTTACATTCCA (SEQ ID NO:38); miR-30b, GCTGAGTGTAGGATGTTTACA (SEQ ID NO:39); miR-30a-s, GCTTCCAGTCGAGGATGTTTACA (SEQ ID NO:40); miR-99b, CGCAAGGTCGGTTCTACGGGTG (SEQ ID NO:41);

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miR-101, TCAGTTATCACAGTACTGTA (SEQ ID NO:42);
miR-122a, ACAAACACCATTGTCACACTCCA (SEQ ID NO:43);
miR-124a, TGGCATTCACCGCGTGCCTTA (SEQ ID NO:44);
miR-125a, CACAGGTTAAAGGGTCTCAGGGA (SEQ ID NO:45);
miR-125b, TCACAAGTTAGGGTCTCAGGGA (SEQ ID NO:46);
miR-127, AGCCAAGCTCAGACGGATCCGA (SEQ ID NO:47);
miR-128, AAAAGAGACCGGTTCACTCTGA (SEQ ID NO:48);
miR-129, GCAAGCCCAGACCGAAAAAAG (SEQ ID NO:49);
miR-130, GCCCTTTTAACATTGCACTC (SEQ ID NO:50);
miR-131, ACTTTCGGTTATCTAGCTTTA (SEQ ID NO:51);
miR-132, ACGACCATGGCTGTAGACTGTTA (SEQ ID NO:52);
miR-143, TGAGCTACAGTGCTTCATCTCA (SEQ ID NO:53).

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Fig. 6. Potential orthologs of lin-4 stRNA. (A) Sequence alignment of *C. elegans* lin-4 stRNA with mouse miR-125a and miR-125b and the *D. melanogaster* miR-125. Differences are highlighted by gray boxes. (B) Northern blot of total RNA isolated from staged populations of *D. melanogaster*, probed for miR-125. E, embryo; L, larval stage; P, pupae; A, adult; S2, Schneider-2 cells.

Fig. 7. Predicted precursor structures of miRNAs, sequence accession numbers and homology information. RNA secondary structure prediction was performed using mfold version 3.1 and manually refined to accommodate G/U wobble base pairs in the helical segments. Dashes were inserted into the secondary structure presentation when asymmetrically bulged nucleotides had to be accommodated. The excised miRNA sequence is underlined. The actual size of the stem-loop structure is not known experimentally and may be slightly shorter or longer than represented. Multicopy miRNAs and their corresponding precursor structures are also shown. In cases where no mouse precursors were yet deposited in the database, the human orthologs are indicated. miRNAs

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which correspond to *D. melanogaster* or human sequences are included. Published *C. elegans* miRNAs [36, 37] are also included in the table. A recent set of new HeLa cell miRNAs is also indicated [46]. If several ESTs were retrieved for one organism in the database, only those with different precursor sequences are listed. miRNA homologs found in other species are indicated. Chromosomal location and sequence accession numbers, and clusters of miRNA genes are indicated. Sequences from cloned miRNAs were searched against mouse and human in GenBank (including trace data), and against *Fugu rubripes* and *Danio rerio* at www.jgi.doe.gov and www.sanger.ac.uk, respectively.

# EXAMPLE 1: MicroRNAs from D. melanogaster and human.

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We previously developed a directional cloning procedure to isolate siRNAs after processing of long dsRNAs in Drosophila melanogaster embryo lysate (8). Briefly, 5' and 3' adapter molecules were ligated to the ends of a size-fractionated RNA population, followed by reverse transcription, PCR amplification, concatamerization, cloning and sequencing. This method, originally intended to isolate siRNAs, led to the simultaneous identification of 14 novel 20- to 23-nt short RNAs which are encoded in the D. melanogaster genome and which are expressed in 0 to 2 h embryos (Table 1). The method was adapted to clone RNAs in a similar size range from HeLa cell total RNA (14), which led to the identification of 19 novel human stRNAs (Table 2), thus providing further evidence for the existence of a large class of small RNAs with potential regulatory roles. According to their small size, we refer to these novel RNAs as microRNAs or miRNAs. The miRNAs are abbreviated as miR-1 to miR-33, and the genes encoding miRNAs are named mir-1 to mir-33. Highly homologous miRNAs are classified by adding a lowercase letter, followed by a dash and a number for designating multiple genomic copies of a mir gene.

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The expression and size of the cloned, endogenous short RNAs was also examined by Northern blotting (Fig. 1, Table 1 and 2). Total RNA isolation was performed by acid guanidinium thiocyanate-phenol-chloroform extraction [45]. Northern analysis was performed as described [1], except that the total RNA was resolved on a 15% denaturing polyacrylamide gel, transferred onto Hybond-N+membrane (Amersham Pharmacia Biotech), and the hybridization and wash steps were performed at 50°C. Oligodeoxynucleotides used as Northern probes were 5′-32P-phosphorylated, complementary to the miRNA sequence and 20 to 25 nt in length.

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5S rRNA was detected by ethidium staining of polyacrylamide gels prior to transfer. Blots were stripped by boiling in 0.1% aqueous sodium dodecylsulfate/0.1x SSC (15 mM sodium chloride, 1.5 mM sodium citrate, pH 7.0) for 10 min, and were re-probed up to 4 times until the 21-nt signals became too weak for detection. Finally, blots were probed for val-tRNA as size marker.

For analysis of D. melanogaster RNAs, total RNA was prepared from different developmental stages, as well as cultured Schneider-2 (S2) cells, which originally derive from 20-24 h D. melanogaster embryos [15] (Fig. 1, Table 1). miR-3 to miR-7 are expressed only during embryogenesis and not at later developmental stages. The temporal expression of miR-1, miR-2 and miR-8 to miR-13 was less restricted. These miRNAs were observed at all developmental stages though significant variations in the expression levels were sometimes observed. Interestingly, miR-1, miR-3 to miR-6, and miR-8 to miR-11 were completely absent from cultured Schneider-2 (S2) cells, which were originally derived from 20-24 h D. melanogaster embryos [15], while miR-2, miR-7, miR-12, and miR-13 were present in S2 cells, therefore indicating cell type-specific miRNA expression. miR-1, miR-8, and miR-12 expression patterns are similar to those of lin-4 stRNA in C. elegans, as their expression is strongly upregulated in larvae and sustained

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to adulthood [16]. miR-9 and miR-11 are present at all stages but are strongly reduced in the adult which may reflect a maternal contribution from germ cells or expression in one sex only.

The mir-3 to mir-6 genes are clustered (Fig. 2A), and mir-6 is present as triple repeat with slight variations in the mir-6 precursor sequence but not in the miRNA sequence itself. The expression profiles of miR-3 to miR-6 are highly similar (Table 1), which suggests that a single embryo-specific precursor transcript may give rise to the different miRNAs, or that the same enhancer regulates miRNA-specific promoters. Several other fly miRNAs are also found in gene clusters (Fig. 2A).

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The expression of HeLa cell miR-15 to miR-33 was examined by Northern blotting using HeLa cell total RNA, in addition to total RNA prepared from mouse kidneys, adult zebrafish, Xenopus laevis ovary, and D. melanogaster S2 cells (Fig. 1B, Table 2). miR-15 and miR-16 are encoded in a gene cluster (Fig. 2B) and are detected in mouse kidney, fish, and very weakly in frog ovary, which may result from miRNA expression in somatic ovary tissue rather than oocytes. mir-17 to mir-20 are also clustered (Fig. 2B), and are expressed in HeLa cells and fish, but undetectable in mouse kidney and frog ovary (Fig. 1, Table 2), and therefore represent a likely case of tissue-specific miRNA expression.

The majority of vertebrate and invertebrate miRNAs identified in this study are not related by sequence, but a few exceptions, similar to the highly conserved let-7 RNA [6], do exist. Sequence analysis of the D. melanogaster miRNAs revealed four such examples of sequence conservation between invertebrates and vertebrates. miR-1 homologs are encoded in the genomes of C. elegans, C. briggsae, and humans, and are found in cDNAs from zebrafish, mouse, cow and human. The expression of mir-1 was detected by Northern blotting in total RNA from adult zebrafish and C. elegans, but not in total RNA from HeLa cells or mouse kidney

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(Table 2 and data not shown). Interestingly, while mir-1 and let-7 are expressed both in adult flies (Fig. 1A) [6] and are both undetected in S2 cells, miR-1 is, in contrast to let-7, undetectable in HeLa cells. This represents another case of tissue-specific expression of a miRNA, and indicates that miRNAs may not only play a regulatory role in developmental timing, but also in tissue specification. miR-7 homologs were found by database searches in mouse and human genomic and expressed sequence tag sequences (ESTs). Two mammalian miR-7 variants are predicted by sequence analysis in mouse and human, and were detected by Northern blotting in HeLa cells and fish, but not in mouse kidney (Table 2). Similarly, we identified mouse and human miR-9 and miR-10 homologs by database searches but only detected mir-10 expression in mouse kidney.

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The identification of evolutionary related miRNAs, which have already acquired multiple sequence mutations, was not possible by standard bioinformatic searches. Direct comparison of the D. melanogaster miRNAs with the human miRNAs identified an 11-nt segment shared between D. melanogaster miR-6 and HeLa miR-27, but no further relationships were detected. One may speculate that most miRNAs only act on a single target and therefore allow for rapid evolution by covariation, and that highly conserved miRNAs act on more than one target sequence, and therefore have a reduced probability for evolutionary drift by covariation [6]. An alternative interpretation is that the sets of miRNAs from D. melanogaster and humans are fairly incomplete and that many more miRNAs remain to be discovered, which will provide the missing evolutionary links.

lin-4 and let-7 stRNAs were predicted to be excised from longer transcripts that contain approximately 30 base-pair stem-loop structures [1, 6]. Database searches for newly identified miRNAs revealed that all miRNAs are flanked by sequences that have the potential to form stable stem-loop structures (Fig. 3 and 4). In many cases, we were able to detect the predicted, approximately 70-nt precursors by Northern blotting (Fig. 1).

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Some miRNA precursor sequences were also identified in mammalian cDNA (EST) databases [27], indicating that primary transcripts longer than 70-nt stem-loop precursors do also exist. We never cloned a 22-nt RNA complementary to any of the newly identified miRNAs, and it is as yet unknown how the cellular processing machinery distinguishes between the miRNA and its complementary strand. Comparative analysis of the precursor stem-loop structures indicates that the loops adjacent to the base-paired miRNA segment can be located on either side of the miRNA sequence (Fig. 3 and 4), suggesting that the 5 or 3 location of the stemclosing loop is not the determinant of miRNA excision. It is also unlikely that the structure, length or stability of the precursor stem is the critical determinant as the base-paired structures are frequently imperfect and interspersed by less stable, non-Watson-Crick base pairs such as G/A, U/U, C/U, A/A, and G/U wobbles. Therefore, a sequence-specific recognition process is a likely determinant for miRNA excision, perhaps mediated by members of the Argonaute (rde-1/ago1/piwi) protein family. Two members of this family, alg-1 and alg-2, have recently been shown to be critical for stRNA processing in C. elegans [13]. Members of the Argonaute protein family are also involved in RNAi and PTGS. In D. melanogaster, these include argonaute2, a component of the siRNA-endonuclease complex (RISC) [17], and its relative aubergine, which is important for silencing of repeat genes [18]. In other species, these include rde-1, argonaute1, and qde-2, in C. elegans [19], Arabidopsis thaliana [20], and Neurospora crassa [21], respectively. The Argonaute protein family therefore represents, besides the RNase III Dicer [12, 13], another evolutionary link between RNAi and miRNA maturation.

Despite advanced genome projects, computer-assisted detection of genes encoding functional RNAs remains problematic [22]. Cloning of expressed, short functional RNAs, similar to EST approaches (RNomics), is a powerful alternative and probably the most efficient method for identification of such novel gene products [23-26]. The number of functional RNAs has been

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widely underestimated and is expected to grow rapidly because of the development of new functional RNA cloning methodologies.

The challenge for the future is to define the function and the potential targets of these novel miRNAs by using bioinformatics as well as genetics, and to establish a complete catalogue of time- and tissue-specific distribution of the already identified and yet to be uncovered miRNAs. lin-4 and let-7 stRNAs negatively regulate the expression of proteins encoded by mRNAs whose 3' untranslated regions contain sites of complementarity to the stRNA [3-5].

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Thus, a series of 33 novel genes, coding for 19- to 23-nucleotide microRNAs (miRNAs), has been cloned from fly embryos and human cells. Some of these miRNAs are highly conserved between vertebrates and invertebrates and are developmentally or tissue-specifically expressed. Two of the characterized human miRNAs may function as tumor suppressors in B-cell chronic lymphocytic leukemia. miRNAs are related to a small class of previously described 21- and 22-nt RNAs (lin-4 and let-7 RNAs), so-called small temporal RNAs (stRNAs), and regulate developmental timing in C. elegans and other species. Similar to stRNAs, miRNAs are presumed to regulate translation of specific target mRNAs by binding to partially complementary sites, which are present in their 3'-untranslated regions.

Deregulation of miRNA expression may be a cause of human disease, and detection of expression of miRNAs may become useful as a diagnostic. Regulated expression of miRNAs in cells or tissue devoid of particular miRNAs may be useful for tissue engineering, and delivery or transgenic expression of miRNAs may be useful for therapeutic intervention. miRNAs may also represent valuable drug targets itself. Finally, miRNAs and their precursor sequences may be engineered to recognize therapeutic valuable targets.

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# EXAMPLE 2: miRNAs from mouse.

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To gain more detailed insights into the distribution and function of miRNAs in mammals, we investigated the tissue-specific distribution of miRNAs in adult mouse. Cloning of miRNAs from specific tissues was preferred over whole organism-based cloning because low-abundance miRNAs that normally go undetected by Northern blot analysis are identified clonally. Also, in situ hybridization techniques for detecting 21-nt RNAs have not yet been developed. Therefore, 19- to 25-nucleotide RNAs were cloned and sequenced from total RNA, which was isolated from 18.5 weeks old BL6 mice. Cloning of miRNAs was performed as follows: 0.2 to 1 mg of total RNA was separated on a 15% denaturing polyacrylamide gel and RNA of 19- to 25-nt size was recovered. A 5´-phosphorylated 3´-adapter oligonucleotide (5 '-pUUUaaccgcgaattccagx: uppercase, RNA; lowercase, DNA; p, phosphate; x, 3'-Amino-Modifier C-7, ChemGenes, Ashland, Ma, USA, Cat. No. NSS-1004; SEQ ID NO:54) and a 5 '-adapter oligonucleotide (5 '-acggaattcctcactAAA: uppercase, RNA; lowercase, DNA; SEQ ID NO:55) were ligated to the short RNAs. RT/PCR was performed with 3'primer (5 '-GACTAGCTGGAATTCGCGGTTAAA; SEQ ID NO:56) and 5 'primer (5 '-CAGCCAACGGAATTCCTCACTAAA; SEQ ID NO:57). In order to introduce Ban I restriction sites, a second PCR was performed using the primer pair 5'-CAGCCAACAGGCACCGAATTCCTCACTAAA (SEQ ID NO:57) and 5'-GACTAGCTTGGTGCCGAATTCGCGGTTAAA (SEQ ID NO:56), followed by concatamerization after Ban I digestion and T4 DNA ligation. Concatamers of 400 to 600 basepairs were cut out from 1.5% agarose gels and recovered by Biotrap (Schleicher & Schuell) electroelution (1x TAE buffer) and by ethanol precipitation. Subsequently, the 3 'ends of the concatamers were filled in by incubating for 15 min at 72°C with Taq polymerase in standard PCR reaction mixture. This solution was diluted 3fold with water and directly used for ligation into pCR2.1 TOPO vectors. Clones were screened for inserts by PCR and 30 to 50 samples were subjected to sequencing. Because RNA was prepared from combining

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tissues of several mice, minor sequence variations that were detected multiple times in multiple clones may reflect polymorphisms rather than RT/PCR mutations. Public database searching was used to identify the genomic sequences encoding the approx. 21-nt RNAs. The occurrence of a 20 to 30 basepair fold-back structure involving the immediate upstream or downstream flanking sequences was used to assign miRNAs [36-38].

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We examined 9 different mouse tissues and identified 34 novel miRNAs, some of which are highly tissue-specifically expressed (Table 3 and Figure 5). Furthermore, we identified 33 new miRNAs from different mouse tissues and also from human Soas-2 osteosarcoma cells (Table 4). miR-1 was previously shown by Northern analysis to be strongly expressed in adult heart, but not in brain, liver, kidney, lung or colon [37]. Here we show that miR-1 accounts for 45% of all mouse miRNAs found in heart, yet miR-1 was still expressed at a low level in liver and midbrain even though it remained undetectable by Northern analysis. Three copies or polymorphic alleles of miR-1 were found in mice. The conservation of tissue-specific miR-1 expression between mouse and human provides additional evidence for a conserved regulatory role of this miRNA. In liver, variants of miR-122 account for 72% of all cloned miRNAs and miR-122 was undetected in all other tissues analyzed. In spleen, miR-143 appeared to be most abundant, at a frequency of approx. 30%. In colon, miR-142-as, was cloned several times and also appeared at a frequency of 30%. In small intestine, too few miRNA sequences were obtained to permit statistical analysis. This was due to strong RNase activity in this tissue, which caused significant breakdown of abundant non-coding RNAs, e.g. rRNA, so that the fraction of miRNA in the cloned sequences was very low. For the same reason, no miRNA sequences were obtained from pancreas.

To gain insights in neural tissue miRNA distribution, we analyzed cortex, cerebellum and midbrain. Similar to heart, liver and small intestine, variants

of a particular miRNA, miR-124, dominated and accounted for 25 to 48% of all brain miRNAs. miR-101, -127, -128, -131, and -132, also cloned from brain tissues, were further analyzed by Northern blotting and shown to be predominantly brain-specific. Northern blot analysis was performed as described in Example 1. tRNAs and 5S rRNA were detected by ethidium staining of polyacrylamide gels prior to transfer to verify equal loading. Blots were stripped by boiling in deionized water for 5 min, and reprobed up to 4 times until the 21-nt signals became too weak for detection.

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miR-125a and miR-125b are very similar to the sequence of C. elegans lin-4 stRNA and may represent its orthologs (Fig. 6A). This is of great interest because, unlike let-7 that was readily detected in other species, lin-4 has acquired a few mutations in the central region and thus escaped bioinformatic database searches. Using the mouse sequence miR-125b, we could readily identify its ortholog in the D. melanogaster genome. miR-125a and miR-125b differ only by a central diuridine insertion and a U to C change. miR-125b is very similar to lin-4 stRNA with the differences located only in the central region, which is presumed to be bulged out during target mRNA recognition [41]. miR-125a and miR-125b were cloned from brain tissue, but expression was also detected by Northern analysis in other tissues, consistent with the role for lin-4 in regulating neuronal remodeling by controlling lin-14 expression [43]. Unfortunately, orthologs to C. elegans lin-14 have not been described and miR-125 targets remain to be identified in D. melanogaster or mammals. Finally, miR-125b expression is also developmentally regulated and only detectable in pupae and adult but not in embryo or larvae of D. melanogaster (Fig. 6B).

Sequence comparison of mouse miRNAs with previously described miRNA reveals that miR-99b and miR-99a are similar to *D. melanogaster*, mouse and human miR-10 as well as *C. elegans* miR-51 [36], miR-141 is similar to *D. melanogaster* miR-8, miR-29b is similar to *C. elegans* miR-83, and miR-131 and miR-142-s are similar to *D. melanogaster* miR-4 and *C.* 

elegans miR-79 [36]. miR-124a is conserved between invertebrates and vertebrates. In this respect it should be noted that for almost every miRNA cloned from mouse was also encoded in the human genome, and frequently detected in other vertebrates, such as the pufferfish, Fugu rubripes, and the zebrafish, Danio rerio. Sequence conservation may point to conservation in function of these miRNAs. Comprehensive information about orthologous sequences is listed in Fig. 7.

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In two cases both strands of miRNA precursors were cloned (Table 3), which was previously observed once for a *C. elegans* miRNA [36]. It is thought that the most frequently cloned strand of a miRNA precursor represents the functional miRNA, which is miR-30c-s and miR-142-as, s and as indicating the 5 ° or 3 ° side of the fold-back structure, respectively.

The mir-142 gene is located on chromosome 17, but was also found at the breakpoint junction of a t(8;17) translocation, which causes an aggressive B-cell leukemia due to strong up-regulation of a translocated MYC gene [44]. The translocated MYC gene, which was also truncated at the first exon, was located only 4-nt downstream of the 3´-end of the miR-142 precursor. This suggests that translocated MYC was under the control of the upstream miR-142 promoter. Alignment of mouse and human miR-142 containing EST sequences indicate an approximately 20 nt conserved sequence element downstream of the mir-142 hairpin. This element was lost in the translocation. It is conceivable that the absence of the conserved downstream sequence element in the putative miR-142/mRNA fusion prevented the recognition of the transcript as a miRNA precursor and therefore may have caused accumulation of fusion transcripts and overexpression of MYC.

miR-155, which was cloned from colon, is excised from the known noncoding BIC RNA [47]. BIC was originally identified as a gene transcriptionally activated by promoter insertion at a common retroviral

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integration site in B cell lymphomas induced by avian leukosis virus. Comparison of BIC cDNAs from human, mouse and chicken revealed 78% identity over 138 nucleotides [47]. The identity region covers the miR-155 fold-back precursor and a few conserved boxes downstream of the fold-back sequence. The relatively high level of expression of BIC in lymphoid organs and cells in human, mouse and chicken implies an evolutionary conserved function, but BIC RNA has also been detected at low levels in non-hematopoietic tissues [47].

Another interesting observation was that segments of perfect complementarity to miRNAs are not observed in mRNA sequences or in genomic sequences outside the miRNA inverted repeat. Although this could be fortuitous, based on the link between RNAi and miRNA processing [11, 13, 43] it may be speculated that miRNAs retain the potential to cleave perfectly complementary target RNAs. Because translational control without target degradation could provide more flexibility it may be preferred over mRNA degradation.

In summary, 63 novel miRNAs were identified from mouse and 4 novel miRNAs were identified from human Soas-2 osteosarcoma cells (Table 3 and Table 4), which are conserved in human and often also in other non-mammalian vertebrates. A few of these miRNAs appear to be extremely tissue-specific, suggesting a critical role for some miRNAs in tissue-specification and cell lineage decisions. We may have also identified the fruitfly and mammalian ortholog of *C. elegans* lin-4 stRNA. The establishment of a comprehensive list of miRNA sequences will be instrumental for bioinformatic approaches that make use of completed genomes and the power of phylogenetic comparison in order to identify miRNA-regulated target mRNAs.

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## Table 1

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D. melanogaster miRNAs. The sequences given represent the most abundant, and typically longest miRNA sequence identified by cloning; miRNAs frequently vary in length by one or two nucleotides at their 3′ termini. From 222 short RNAs sequenced, 69 (31%) corresponded to miRNAs, 103 (46%) to already characterized functional RNAs (rRNA, 7SL RNA, tRNAs), 30 (14%) to transposon RNA fragments, and 20 (10%) sequences with no database entry. The frequency (freq.) for cloning a particular miRNA relative to all identified miRNAs is indicated in percent. Results of Northern blotting of total RNA isolated from staged populations of D. melanogaster are summarized. E, embryo; L, larval stage; P, pupae; A, adult; S2, Schneider-2 cells. The strength of the signal within each blot is represented from strongest (+ + +) to undetected (-). let-7 stRNA was probed as control. Genbank accession numbers and homologs of miRNAs identified by database searching in other species are provided as supplementary material.

	miRNA	sequence (5' to 3')	freq.	E	E	L1+	L3	P	Α	S2
			(%)	0-3 h	0-6 h	L2				
	miR-1	UGGAAUGUAAAGAAGUAUGGAG	32	+	+	++	++	++	++	-
,		(SEQ ID NO:58)				+	+		+	
20	miR-2a*	UAUCACAGCCAGCUUUGAUGAGC	3					1	<b></b>	
		(SEQ ID NO:59)								
	miR-2b*	UAUCACAGCCAGCUUUGAGGAGC	3	++	++	++	++	++	+	++
		(SEQ ID NO:60)					+			+
	miR-3	UCACUGGGCAAAGUGUGUCUCA#	9	+++	+++	-	-	-	-	-
25	miR-4	AUAAAGCUAGACCAUUGA (SEQ ID NO:62)	6	+++	+++	-	-	-	-	-
	miR-5	AAAGGAACGAUCGUUGUGAUAUG (SEQ ID NO:63)	1	+++	+++	+/-	+/-	-	-	-
ŀ	miR-6	UAUCACAGUGGCUGUUCUUUU	13	+++	+++	+/-				
	11111 \(\frac{1}{2}\)	(SEO ID NO:64)	13	777	777	+/-	+/-	-	-	-
ŀ	miR-7	UGGAAGACUAGUGAUUUUGUUGU	4	+++	++	+/-	+/-	+/-	+/-	+/
		(SEQ ID NO:65)	·			·	,		''	
Ì	miR-8	UAAUACUGUCAGGUAAAGAUGUC	3	+/-	+/-	++	++	+	++	-
		(SEQ ID NO:66)				+	+		+	

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miR-9	UCUUUGGUUAUCUAGCUGUAUGA	7	+++	++	++	++	++	+/-	Τ-
	(SEQ ID NO:67)				+	+	+		
miR-10	ACCCUGUAGAUCCGAAUUUGU	1	+	+	++	++	+/-	+	<del> </del>
	(SEQ ID NO:68)					+			
miR-11	CAUCACAGUCUGAGUUCUUGC	7	+++	+++	++	++	++	+	-
	(SEQ ID NO:69)	٠.			+	+	+	· ·	
miR-12	UGAGUAUUACAUCAGGUACUGGU	7	+	+	++	++	+	++	+/-
-	(SEQ ID NO:70)							+	
miR-13a*	UAUCACAGCCAUUUUGACGAGU	1	+++	+++	++	++	+	++	++
	(SEQ ID NO:71)				+	+		+ .	+
miR-13b*	UAUCACAGCCAUUUUGAUGAGU	Ó		•	1			:	<u> </u>
	(SEQ ID NO:72)								
miR-14	UCAGUCUUUUUCUCUCUCCUA	1 ·		- '	1	-	† <del>-</del>		-
:	(SEQ ID NO:73)								
let-7	UGAGGUAGUAGGUUGUAUAGUU	0	-	T -	T=	-	++	++	-
	(SEQ ID NO:74)			1			+	_	

10 # = (SEQ ID NO:61)

<sup>\*</sup>Similar miRNA sequences are difficult to distinguish by Northern blotting because of potential cross-hybridization of probes.

Table 2

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Human miRNAs. From 220 short RNAs sequenced, 100 (45%) corresponded to miRNAs, 53 (24%) to already characterized functional RNAs (rRNA, snRNAs, tRNAs), and 67 (30%) sequences with no database entry. Results of Northern blotting of total RNA isolated from different vertebrate species and S2 cells are indicated. For legend, see Table 1.

	miRNA	sequence (5' to 3')	freq.	HeLa	. mouse	adult -	frog ·	S2 :
,			(%)	cells	, kidney	fish	ovary	
	let-7a*	UGAGGUAGUAGGUUGUAUAGUU#	10 -	+++	+++	+++	_	_
10	let-7b*	UGAGGUAGUAGGUUGUGGUU	: 13					·
<i>"</i> .		(SEQ ID NO:76)		٠. ٠		San Comment	***	
	let-7c*	UGAGGUAGUAGGUÜ	3					
		(SEQ ID NO:77)						
İ	let-7d*	AGAGGUAGUAGGUUGCAUAGU	2	+++	+++	+++	-	-
		(SEQ ID NO:78)		,				
	let-7e*	UGAGGUAGGAGGUUGUAUAGU	2	+++	+++	+++	-	-
į		(SEQ ID NO:79)						
	let-7f*	UGAGGUAGUAGAUUGUAUAGUU	1					
		(SEQ ID NO:80)						
15	miR-15	UAGCAGCACAUAAUGGUUUGUG	3	+++	++	+	+/-	-
		(SEQ ID NO:81)						
	miR-16	UAGCAGCACGUAAAUAUUGGCG	10	+++	+	+/-	+/-	
		(SEQ ID NO:82)						
ł	miR-17	ACUGCAGUGAAGGCACUUGU	1	+++	-	-	-	-
		(SEQ ID NO:83)						
	miR-18	UAAGGUGCAUCUAGUGCAGAUA	2	+++	-	-	-	-
		(SEQ ID NO:84)						
	miR-19a*	UGUGCAAAUCUAUGCAAAACUGA	1	+++	-	+/-	-	-
	-	(SEQ ID NO:85)						
20	miR-19b*	UGUGCAAAUCCAUGCAAAACUGA	3					
		(SEQ ID NO:86)						
	miR-20	UAAAGUGCUUAUAGUGCAGGUA	4	+++	-	+	-	-
		(SEQ ID NO:87)						
	miR-21	UAGCUUAUCAGACUGAUGUUGA	10	+++	÷	++	-	-
		(SEQ ID NO:88)						
	miR-22	AAGCUGCCAGUUGAAGAACUGU	10	+++	+++	+	+/-	-
		(SEQ ID NO:89)						
	miR-23	AUCACAUUGCCAGGGAUUUCC	2	+++	+++	+++	+	-
		(SEQ ID NO:90)						
		war						

ſ	miR-24	UGGCUCAGUUCAGCAGGAACAG	4	++	+++	++	T =	l <u> </u>
		(SEQ ID NO:91)						
ł	miR-25	CAUUGCACUUGUCUCGGUCUGA	3	+++	+	++	<del> </del>	_
		(SEQ ID NO:92)						
ŀ	miR-26a*	UUCAAGUAAUCCAGGAUAGGCU	2	+	++	+++	-	-
		(SEQ ID NO:93)						
Ì	miR-26b*	UUCAAGUAAUUCAGGAUAGGUU	1			<u> </u>		-
	•	(SEQ ID NO:94)						
5	miR-27	UUCACAGUGGCUAAGUUCCGCU	2	+++	+++	.++.	-	-
		(SEQ ID NO:95)				,		
	miR-28	AAGGAGCUCACAGUCUAUUGAG	2	7+++	+++	-	-	
		(SEQ ID NO:96)						
	miR-29	CUAGCACCAUCUGAAAUCGGUU	2	+	+++	+/-	-	-
		(SEQ ID NO:97)	s.'	., .,				, -
	miR-30	CUUUCAGUCGGAUGUUUGCAGC ·	2	+++	+++ :	+++	- 2.00	-
		(SEQ ID NO:98)						
	miR-31	GGCAAGAUGCUGGCAUAGCUG	2	+++	-	-	-	-
		(SEQ ID NO:99)					İ	
10	miR-32	UAUUGCACAUUACUAAGUUGC	1 .	-	-		-	-
		(SEQ ID NO:100)						
Ì	miR-33	GUGCAUUGUAGUUGCAUUG	1	-	-	-	-	-
		(SEQ ID NO:101)						
	miR-1	UGGAAUGUAAAGAAGUAUGGAG	0	-	-	.+	-	-
		(SEQ ID NO:102)						
	miR-7	UGGAAGACUAGUGAUUUUGUUGU	0	+	+-	+/-	-	+/-
		(SEQ ID NO:103)						
j	miR-9	UCUUUGGUUAUCUAGCUGUAUGA	0	-	-	-	-	-
		(SEQ ID NO:104)						
15	miR-10	ACCCUGUAGAUCCGAAUUUGU	0	-	+	-	-	-
		(SEQ ID NO:105)				1	]	

# = (SEQ ID NO:75)

<sup>\*</sup>Similar miRNA sequences are difficult to distinguish by Northern blotting because of potential cross-hybridization of probes.

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Table 3

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Mouse miRNAs. The sequences indicated represent the longest miRNA sequences identified by cloning. The 3′-terminus of miRNAs is often truncated by one or two nucleotides. miRNAs that are more than 85% identical in sequence (i.e. share 18 out of 21 nucleotides) or contain 1- or 2-nucleotide internal deletions are referred to by the same gene number followed by a lowercase letter. Minor sequence variations between related miRNAs are generally found near the ends of the miRNA sequence and are thought to not compromise target RNA recognition. Minor sequence variations may also represent A to G and C to U changes, which are accommodated as G-U wobble base pairs during target recognition. miRNAs with the suffix -s or -as indicate RNAs derived from either the 5′-half or the 3′-half of a miRNA precursor. Mouse brains were dissected into midbrain, mb, cortex, cx, cerebellum, cb. The tissues analyzed were heart, ht; liver, lv; small intestine, si; colon, co; cortex, ct; cerebellum, cb; midbrain, mb.

	miRNA	sequence (5' to 3')			Numb	er o	f clo	ones		
20			ht	lv	sp	si	со	cx	cb	mb
	let-7a	UGAGGUAGUAGGUUGUAUAGUU (SEQ ID NO:106)		3			1	1		7
	let-7b	UGAGGUAGUAGGUUGUGGGUU (SEQ ID NO:107)		1	1				2	5
	let-7c	UGAGGUAGUAGGUUGUAUGGUU (SEQ ID NO:108)		2				2	5	19
	let-7d	AGAGGUAGUAGGUUGCAUAGU (SEQ ID NO:109)	2				2	2		2
25	1et-7e	UGAGGUAGGAGGUUGUAUAGU (SEQ ID NO:110)		-	1					2
	let-7f	UGAGGUAGUAGAUUGUAUAGUU (SEQ ID NO:111)			2				3	3
	let-7g	UGAGGUAGUAGUUUGUACAGUA (SEQ ID NO:112)						1	1	2
	let-7h	UGAGGUAGUAGUGUACAGUU (SEQ ID NO:113)						1	1	

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	let-7i	UGAGGUAGUUUGUGCU (SEQ ID NO:114)						1	1	
	miR-1b	UGGAAUGUAAAGAAGUAUGUAA (SEQ ID NO:115)	4	2						1
	miR-1c	UGGAAUGUAAAGAAGUAUGUAC (SEQ ID NO:116)	7							
	miR-1d	UGGAAUGUAAAGAAGUAUGUAUU (SEQ ID NO:117)	16							1
5	miR-9	UCUUUGGUUAUCUAGCUGUAUGA (SEQ ID NO:118)					•	3	4	4
	miR-15a	UAGCAGCACAUAAUGGUUUGUG (SEQ ID NO:119)	1 .							2
	miR-15b	UAGCAGCACAUCAUGGUUUACA (SEQ ID NO:120)	1 .							
ę i	miR-16	UAGCAGCACGUAAAUAUUGGCG (SEQ ID NO:121)	1 .			1	2-	1	2	3
	miR-18	UAAGGUGCAUCUAGUGCAGAUA (SEQ ID NO:122)			1		·			
10	miR-19b	UGUGCAAAUCCAUGCAAAACUGA (SEQ ID NO:123)			1					
	miR-20	UAAAGUGCUUAUAGUGCAGGUAG (SEQ ID NO:124)					1			
	miR-21 :	UAGCUUAUCAGACUGAUGUUGA (SEQ ID NO:125)	1		1 .	2	1			
	miR-22	AAGCUGCCAGUUGAAGAACUGU (SEQ ID NO:126)	2	1		1			1	2
	miR-23a	AUCACAUUGCCAGGGAUUUCC (SEQ ID NO:127)	1							
15	miR-23b	AUCACAUUGCCAGGGAUUACCAC (SEQ ID NO:128)						1		
	miR-24	UGGCUCAGUUCAGCAGGAACAG (SEQ ID NO:129)	1				1	1		1
	miR-26a	UUCAAGUAAUCCAGGAUAGGCU (SEQ ID NO:130)							3	2
	miR-26b	UUCAAGUAAUUCAGGAUAGGUU (SEQ ID NO:131)		2				4	1	
	miR-27a	UUCACAGUGGCUAAGUUCCGCU (SEQ ID NO:132)	1		2		1	1	2	1
20	miR-27b	UUCACAGUGGCUAAGUUCUG (SEQ ID NO:133)								1
	miR-29a	CUAGCACCAUCUGAAAUCGGUU (SEQ ID NO:134)	1				1		1	
	miR-29b/miR-102	UAGCACCAUUUGAAAUCAGUGUU (SEQ ID NO:135)	1				1	5		3
	miR-29c/	UAGCACCAUUUGAAAUCGGUUA (SEQ ID NO:136)	1					3		1

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	miR-30a-s/miR-97	UGUAAACAUCCUCGACUGGAAGC (SEQ ID NO:137)				1	1		1
	miR-30a-asª	CUUUCAGUCGGAUGUUUGCAGC (SEQ ID NO:138)						1	
	miR-30b	UGUAAACAUCCUACACUCAGC (SEQ ID NO:139)				1		2	
	miR-30c	UGUAAACAUCCUACACUCUCAGC (SEQ ID NO:140)	2				1	1	
5	miR-30d	UGUAAACAUCCCCGACUGGAAG (SEQ ID NO:141)			1	·			
	miR-99a/miR-99	ACCCGUAGAUCCGAUCUUGU (SEQ ID NO:142)				. •	1		
	miR-99b	CACCCGUAGAACCGACCUUGCG (SEQ ID NO:143)		4				1	
	miR-101	UACAGUACUGUGAŲAACUGA (SEQ ID NO:144)			•		2	1	1
-,	miR-122a	UGGAGUGUGACAAUGGUGUUUGU (SEQ ID NO:145)			3				
10	miR-122b	UGGAGUGUGACAAUGGUGUUUGA (SEQ ID NO:146)			11				
	miR-122a,b	UGGAGUGUGACAAUGGUGUUUG (SEQ ID NO:147)			23				
	miR-123	CAUUAUUACUUUUGGUACGCG (SEQ ID NO:148)	. 1		2				
	miR-124a <sup>b</sup>	UUAAGGCACGCGG-UGAAUGCCA (SEQ ID NO:149)				1	37	41	24
	miR-124b	UUAAGGCACGCGGGUGAAUGC (SEQ ID NO:150)					1	3	
15	miR-125a	UCCCUGAGACCCUUUAACCUGUG (SEQ ID NO:151)					1	1	
	miR-125b	UCCCUGAGACCCUAACUUGUGA (SEQ ID NO:152)					1		
	miR-126	UCGUACCGUGAGUAAUAAUGC (SEQ ID NO:153)	4					1	
	miR-127	UCGGAUCCGUCUGAGCUUGGCU (SEQ ID NO:154)						1	
	miR-128	UCACAGUGAACCGGUCUCUUUU (SEQ ID NO:155)					2	2	2
20	miR-129	CUUUUUUCGGUCUGGGCUUGC (SEQ ID NO:156)						1	
	miR-130	CAGUGCAAUGUUAAAAGGGC (SEQ ID NO:157)						1	
	miR-131	UAAAGCUAGAUAACCGAAAGU (SEQ ID NO:158)					1	1	1
	miR-132	UAACAGUCUACAGCCAUGGUCGU (SEQ ID NO:159)						1	

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			•							
	miR-133	UUGGUCCCCUUCAACCAGCUGU (SEQ ID NO:160)	4					1		
	miR-134	UGUGACUGGUUGACCAGAGGGA (SEQ ID NO:161)						1		
	miR-135	UAUGGCUUUUUAUUCCUAUGUGAA (SEQ ID NO:162)						1		
	miR-136	ACUCCAUUUGUUUUGAUGAUGGA (SEQ ID NO:163)						1		• •
5	miR-137	UAUUGCUUAAGAAUACGCGUAG (SEQ ID NO:164)						1 -		1
	miR-138	AGCUGGUGUUGUGAAUC (SEQ ID NO:165)		,				1		
	miR-139	UCUACAGUGCACGUGUCU (SEQ ID NO:166)					. 1	1		
	miR-140	AGUGGUUUUACCCUAUGGUAG (SEQ ID NO:167)		. *-	*	1			,	er ar
	miR-141	AACACUGUCUGGUAAAGAUGG (SEQ ID NO:168)			1	1		1		
10	miR-142-s	CAUAAAGUAGAAAGCACUAC (SEQ ID NO:169)				1	1			
	miR-142-as <sup>b</sup>	UGUAGUGUUUCCUACUUUAUGG (SEQ ID NO:170)			1	1	6			
	miR-143	UGAGAUGAAGCACUGUAGCUCA (SEQ ID NO:171)	3		7			2		1
	miR-144	UACAGUAUAGAUGAUGUACUAG (SEQ ID NO:172)	2				1			
	miR-145	GUCCAGUUUUCCCAGGAAUCCCUU (SEQ ID NO:173)	1							
15	miR-146	UGAGAACUGAAUUCCAUGGGUUU (SEQ ID NO:174)	1							
	miR-147	GUGUGUGGAAAUGCUUCUGCC (SEQ ID NO:175)			1.					
	miR-148	UCAGUGCACUACAGAACUUUGU (SEQ ID NO:176)			1					
	miR-149	UCUGGCUCCGUGUCUUCACUCC (SEQ ID NO:177)	1							
	miR-150	UCUCCCAACCCUUGUACCAGUGU (SEQ ID NO:178)					1			
20	miR-151	CUAGACUGAGGCUCCUUGAGGU (SEQ ID NO:179)					1			
	miR-152	UCAGUGCAUGACAGAACUUGG (SEQ ID NO:180)					1			
	miR-153	UUGCAUAGUCACAAAAGUGA (SEQ ID NO:181)								1
	miR-154	UAGGUUAUCCGUGUUGCCUUCG (SEQ ID NO.182)								1

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miR-155

UUAAUGCUAAUUGUGAUAGGGG (SEQ ID NO:183)

1

The originally described miR-30 was renamed to miR-30a-as in order to distinguish it from the miRNA derived from the opposite strand of the precursor encoded by the mir-30a gene. miR-30a-s is equivalent to miR-97 [46].

<sup>b</sup>A 1-nt length heterogeneity is found on both 5' and 3' end. The 22-nt miR sequence is shown, but only 21-nt miRNAs were cloned.

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## Table 4

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15

Mouse and human miRNAs. The sequences indicated represent the longest miRNA sequences identified by cloning. The 3' terminus of miRNAs is often truncated by one or two nucleotides. miRNAs that are more than 85% identical in sequence (i.e. share 18 out of 21 nucleotides) or contain 1- or 2-nucleotide internal deletions are referred to by the same gene number followed by a lowercase letter. Minor sequence variations between related miRNAs are generally found near the ends of the miRNA sequence and are thought to not compromise target RNA recognition. Minor sequence variations may also represent A to G and C to U changes; which are accommodated as G-U wobble base pairs during target recognition. Mouse brains were dissected into midbrain, mb, cortex, cx, cerebellum, cb. The tissues analyzed were lung, ln; liver, lv; spleen, sp; kidney, kd; skin, sk; testis, ts; ovary, ov; thymus, thy; eye, ey; cortex, ct; cerebellum, cb; midbrain, mb. The human osteosarcoma cells SAOS-2 cells contained an inducible p53 gene (p53-, uninduced p53; p53+, induced p53); the differences in miRNAs identified from induced and uninduced SAOS cells were not statistically significant.

					mnu	number of clones	lones					
വ	miRNA	Sequence (5' to 3')		٠.								
				-	mouse tissues	nes			huma	human SAOS-		
									7	2 cells		
			ln lv	g.	kd sk	ts ov	, thy	ey	p53-	p53+	£ .	
	miR-C1	AACAUUCAACGCUGUCGGUGAGU	1		-			2			(SEQ ID NO.184)	0.184)
10	miR-C2	UUUGGCAAUGGUAGAACUCACA						-			(SEQ ID NO.185)	0.185)
	miR-C3	UAUGGCACUGGUAGAAUUCACUG						-			(SEQ ID NO.186)	0.186)
	miR-C4	cunnuneceencueecunenn				_		_	_		(SEQ ID NO.187)	0.187)
	miR-C5	UGGACGGAGAACUGAUAAGGGU						7	٠.		(SEQ ID NO.188)	0.188)
	miR-C6	UGGAGAGAAAGGCAGUUC									(SEQ ID NO.189)	0.189)
15	miR-C7	CAAAGAAUUCUCCUUUUGGGCUU						-	-		(SEQ ID NO.190)	0.190)
	miR-C8	UCGUGUCUUGUGUUGCAGCCGG		_							(SEQ ID NO.191)	0.191)
	miR-C9	UAACACUGUCUGGUAACGAUG									(SEQ ID NO.192)	0.192)
	miR-C10	CAUCCCUUGCAUGGUGGAGGGU									(SEQ ID NO.193)	0.193)
	miR-C11	GUGCCUACUGAGCUGACAUCAGU									(SEQ ID NO.194)	0.194)
20	miR-C12	UGAUAUGUUUGAUAUAUUAGGU		7	->						(SEQ ID NO.195)	0.195)
	miR-C13	CAACGGAAUCCCAAAAGCAGCU		2 1							(SEQ ID NO.196)	0.196)
	miR-C14	CUGACCUAUGAAUUGACA	2						•	•	(SEQ ID NO.197)	(191)

(SEQ ID NO.198) (SEQ ID NO.199) (SEQ ID NO.200)	(SEQ ID NO.201) (SEQ ID NO.202) (SEQ ID NO.203)	(SEQ ID NO.205) (SEQ ID NO.207) (SEQ ID NO.207) (SEQ ID NO.208)	(SEQ ID NO.210) (SEQ ID NO.211) (SEQ ID NO.212) (SEQ ID NO.213) (SEQ ID NO.214)	(SEQ ID NO.215) (SEQ ID NO.216) (SEQ ID NO.217)
		<b>-</b>		· .
	<del></del>		1 2 2 1	
	64	7		
UACCACAGGGUAGAACCACGGA AACUGGCCUACAAAGUCCCAG UGUAACAGCAACUCCAUGUGGA	UAGGUAGUUCAUGUUGGC UAGGUAGUUCAUGUUGGG UUCACCACCUUCUCCACCCAGC	CCCAGUGUUCAGACUACCUGUU UAAUACUGCCUGGUAAUGAUGAC UACUCAGUAAGGCAUGGAAGA AGAGGUAUAGGACCAUGGAAGA	UUCCCUUUGUCAUCCUAUGCCUG UCCUUCAUUCCACCGGAGUCUG GUGAAAUGUUAGGACCACUAGA UGGAAUGUAAGGAAGUGUGUGG	CCCUGUAGAACCGAAUUUGUGU , AACCCGUAGAUCCGAACUUGUGAA GCUUCUCCUGGCUCUCCCUC
miR-C15 miR-C16 miR-C17	miR-C18 miR-C19 miR-C20 miR-C21	miR-C23 miR-C24 miR-C25 miR-C25	miR-C27 miR-C28 miR-C29 miR-C30	miR-C32 miR-C33 miR-C34
	വ	10	<del>/-</del>	20

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## Table 5

miRNA

seguence (5' to 3')

*D. melanogaster* miRNA sequences and genomic location. The sequences given represent the most abundant, and typically longest miRNA sequences identified by cloning. It was frequently observed that miRNAs vary in length by one or two nucleotides at their 3´-terminus. From 222 short RNAs sequenced, 69 (31%) corresponded to miRNAs, 103 (46%) to already characterized functional RNAs (rRNA, 7SL RNA, tRNAs), 30 (14%) to transposon RNA fragments, and 20 (10%) sequences with no database entry. RNA sequences with a 5´-guanosine are likely to be underrepresented due to the cloning procedure (8). miRNA homologs found in other species are indicated. Chromosomal location (chr.) and GenBank accession numbers (acc. nb.) are indicated. No ESTs matching miR-1 to miR-14 were detectable by database searching.

chr acc nh

remarks

	miRNA	sequence (5° to 3°)	chr., acc. nb.	remarks
15		•		:
	miR-1	UGGAAUGUAAAGAAGUAUGGAG	2L, AE003667	homologs: <i>C. briggsae</i> , G20U,
	•	(SEQ ID NO:58)		AC87074; C.elegans G20U,
				U97405; mouse, G20U, G22U,
				AC020867; human, chr. 20,
				G20U, G22U, AL449263; ESTs:
				zebrafish, G20U, G22U, BF157-
				601; cow, G20U, G22U, BE722-
				224; human, G20U, G22U,
				Al220268
	miR-2a	UAUCACAGCCAGCUUUGAUGAGC	2L, AE003663	2 precursor variants clustered
		(SEQ ID NO:59)		with a copy of <i>mir-2b</i>
20	miR-2b	UAUCACAGCCAGCUUUGAGGAGC	2L, AE003620	2 precursor variants
		(SEQ ID NO:60)	2L, AE003663	
	miR-3	UCACUGGGCAAAGUGUGUCUCA	2R, AE003795	in cluster <i>mir-3</i> to <i>mir-6</i>
		(SEQ ID NO:61)		
	miR-4	AUAAAGCUAGACAACCAUUGA	2R, AE003795	in cluster mir-3 to mir-6
25		(SEQ ID NO:62)		

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	miR-5	AAAGGAACGAUCGUUGUGAUAUG (SEQ ID NO:63)	2R, AE003795	in cluster <i>mir-</i> 3 to <i>mir-</i> 6
	miR-6	UAUCACAGUGGCUGUUCUUUUU (SEQ ID NO:64)	2R, AE003795	in cluster <i>mir-3</i> to <i>mir-6</i> with 3 variants
5	miR-7	UGGAAGACUAGUGAUUUUGUUGU (SEQ ID NO:65)	2R, AE003791	homologs: human, chr. 19 AC006537, EST BF373391; mouse chr. 17 AC026385, EST AA881786
	miR-8	UAAUACUGUCAGGUAAAGAUGUC (SEQ ID NO:66)	2R, AE003805	
10	miR-9	UCUUUGGUUAUCUAGCUGUAUGA (SEQ ID NO:67)	3L, AE003516	homologs: mouse, chr. 19, AF155142; human, chr. 5, AC026701, chr. 15, AC005316
	miR-10	ACCCUGUAGAUCCGAAUUUGU (SEQ ID NO:68)	AE001574	homologs: mouse, chr 11, AC011194; human, chr. 17, AF287967
	miR-11	CAUCACAGUCUGAGUUCUUGC (SEQ ID NO:69)	3R, AE003735	intronic location
15	miR-12	UGAGUAUUACAUCAGGUACUGGU (SEQ ID NO:70)	X, AE003499	intronic location
÷	miR-13a	UAUCACAGCCAUUUUGACGAGU (SEQ ID NO:71)	3R, AE003708 X, AE003446	<i>mir-13a</i> clustered with <i>mir-13b</i> on chr. 3R
20	miR-13b	UAUCACAGCCAUUUUGAUGAGU (SEQ ID NO:72)	3R, AE003708	<i>mir-13a</i> clustered with <i>mir-13b</i> on chr. 3R
	miR-14	UCAGUCUUUUUCUCUCUCCUA (SEQ ID NO:73)	2R, AE003833	no signal by Northern analysis

Table 6

Human miRNA sequences and genomic location. From 220 short RNAs sequenced, 100 (45%) corresponded to miRNAs, 53 (24%) to already characterized functional RNAs (rRNA, snRNAs, tRNAs), and 67 (30%) sequences with no database entry. For legend, see Table 1.

•	miRNA	sequence (5' to 3')	chr. or EST,	remarks*
			acc. nb.	
	let-7a	UGAGGUAGUAGGUUGUAUAGUU	9, AC007924,	sequences of chr 9 and 17
10		(SEQ ID NO:75)	11, AP001359,	identical and clustered with let-7f,
	•	•	17, AC087784,	homologs: C. elegans, AF274345;
	·.		22, AL049853	C. briggsae, AF210771, D.
				melanogaster, AE003659
	let-7b	UGAGGUAGUAGGUUGUGUGGUU	22, AL049853†,	homologs: mouse, EST Al481799;
	•	(SEQ ID NO:76)	ESTs, Al382133,	rat, EST, BE120662
	•	·	AW028822	
				•
	let-7c	UGAGGUAGUAGGUUGUAUGGUU	21, AP001667	Homologs: mouse, EST,
		(SEQ ID NO:77)		AA575575
15	let-7d	AGAGGUAGUAGGUUGCAUAGU	17, AC087784,	identical precursor sequences
		(SEQ ID NO:78)	9, AC007924	•
				*
	let-7e	UGAGGUAGGAGGUUGUAUAGU	19, AC018755	
		(SEQ ID NO:79)		
	let-7f	UGAGGUAGUAGAUUGUAUAGUU	9, AC007924,	sequences of chr 9 and 17
20		(SEQ ID NO:80)	17, AC087784,	identical and clustered with let-7a
			X, AL592046	
	miR-15	UAGCAGCACAUAAUGGUUUGUG	13, AC069475	in cluster with <i>mir-16</i> homolog
		(SEQ ID NO:81)		
	miR-16	UAGCAGCACGUAAAUAUUGGCG	13, AC069475	in cluster with <i>mir-15</i> homolog
		(SEQ ID NO:82)		

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			- 41 -	
	miR-17	ACUGCAGUGAAGGCACUUGU (SEQ ID NO:83)	13, AL138714	in cluster with <i>mir-17</i> to <i>mir-20</i>
	miR-18	UAAGGUGCAUCUAGUGCAGAUA (SEQ ID NO:84)	13, AL138714	in cluster with <i>mir-17</i> to <i>mir-20</i>
5	miR-19a	UGUGCAAAUCUAUGCAAAACUG A (SEQ ID NO:85)	13, AL138714	in cluster with <i>mir-17</i> to <i>mir-20</i>
	miR-19b	UGUGCAAAUCCAUGCAAAACUG A (SEQ ID NO:86)	13, AL138714, X, AC002407	in cluster with <i>mir-17</i> to <i>mir-20</i>
10	miR-20	UAAAGUGCUUAUAGUGCAGGUA (SEQ ID NO:87)	13, AL138714	in cluster with <i>mir-17</i> to <i>mir-20</i>
•	miR-21	UAGCUUAUCAGACUGAUGUUGA (SEQ ID NO:88)	17, AC004686, EST, BF326048	homologs: mouse, EST, AA209594
	miR-22	AAGCUGCCAGUUGAAGAACUGU (SEQ ID NO:89)	ESTs, AW961681†, AA456477, AI752503, BF030303, HS1242049	human ESTs highly similar; homologs: mouse, ESTs, e.g. AA823029; rat, ESTs, e.g. BF543690
15	miR-23	AUCACAUUGCCAGGGAUUUCC (SEQ ID NO:90)	19, AC020916	homologs: mouse, EST, AW124037;rat, EST, BF402515
	miR-24	UGGCUCAGUUCAGCAGGAACAG (SEQ ID NO:91)	9, AF043896, 19, AC020916	homologs: mouse, ESTs, AA111466, Al286629; pig, EST, BE030976
20	miR-25	CAUUGCACUUGUCUCGGUCUGA (SEQ ID NO:92)	7, AC073842, EST, BE077684	human chr 7 and EST identical; highly similar precursors in mouse ESTs (e.g. Al595464); fish precursor different STS: G46757
	miR-26a	UUCAAGUAAUCCAGGAUAGGCU (SEQ ID NO:93)	3, AP000497	

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	miR-26b	UUCAAGUAAUUCAGGAUAGGUU (SEQ ID NO:94)	2, AC021016	
	miR-27	UUCACAGUGGCUAAGUUCCGCU (SEQ ID NO:95)	19, AC20916	U22C mutation in human genomic sequence
5	miR-28	AAGGAGCUCACAGUCUAUUGAG (SEQ ID NO:96)	3, AC063932	•
	miR-29	CUAGCACCAUCUGAAAUCGGUU (SEQ ID NO:97)	7, AF017104	
10	miR-30	CUUUCAGUCGGAUGUUUGCAGC (SEQ ID NO:98)	6, AL035467	
*	miR-31	GGCAAGAUGCUGGCAUAGCUG (SEQ ID NO:99)	9, AL353732	
	miR-32	UAUUGCACAUUACUAAGUUGC (SEQ ID NO:100)	9, AL354797	not detected by Northern blotting
15	miR-33	GUGCAUUGUAGUUGCAUUG (SEQ ID NO:101)	22, Z99716	not detected by Northern blotting

<sup>\*</sup>If several ESTs were retrieved for one organism in the database, only those with different precursor sequences are listed.

<sup>20 †</sup>precursor structure shown in Fig. 4.

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## Claims

1. Isolated nucleic acid molecule comprising

5

- (a) a nucleotide sequence as shown in Table 1, Table 2, Table 3 or Table 4 or a precursor thereof as shown in Figure 3, Figure 4 or Figure 7.
- 10 (b) a nucleotide sequence which is the complement of (a),
  - (c) a nucleotide sequence which has an identity of at least 80% to a sequence of (a) or (b) and/or
- a nucleotide sequence which hybridizes under stringent conditions to a sequence of (a), (b) and/or (c).
  - 2. The nucleic acid molecule of claim 1, wherein the identity of sequence (c) is at least 90%.

- 3. The nucleic acid molecule of claim 1, wherein the identity of sequence (c) is at least 95%.
- 4. The nucleic acid molecule of any one of claims 1-3, which is selected from miR 1-14 as shown in Table 1 or miR 15-33 as shown in Table 2 or miR 1-155 as shown in Table 3 or miR-C1-34 as shown in Table 4 or a complement thereof.
- 5. The nucleic acid molecule of any one of claims 1-3, which is selected from mir 1-14 as shown in Figure 3 or let 7a-7f or mir 15-33, as shown in Figure 4 or let 7a-i or mir 1-155 or mir-c1-34, as shown in Figure 7 or a complement thereof.

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- 6. The nucleic acid molecule of any one of claims 1-4 which is a miRNA molecule or an analog thereof having a length of from 18-25 nucleotides.
- 7. The nucleic acid molecule of any one of claims 1-3 or 5, which is a miRNA precursor molecule having a length of 60-80 nucleotides or a DNA molecule coding therefor.

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- 8. The nucleic acid molecule of any one of claims 1-7, which is single-stranded.
- 9. The nucleic acid molecule of any one of claims 1-7, which is at least partially double-stranded.
- 10. The nucleic acid molecule of any one of claims 1-9, which is selected from RNA, DNA or nucleic acid analog molecules.
  - 11. The nucleic acid molecule of claim 10, which is a molecule containing at least one modified nucleotide analog.
- 20 12. The nucleic molecule of claim 10 which is a recombinant expression vector.
- 13. A pharmaceutical composition containing as an active agent at least one nucleic acid molecule of any one of claims 1-12 and optionally a pharmaceutically acceptable carrier.
  - 14. The composition of claim 13 for diagnostic applications.
  - 15. The composition of claim 13 for therapeutic applications.
  - 16. The composition of any one of claims 13-15 as a marker or a modulator for developmental or pathogenic processes.

17. The composition of claim 13 as a marker or modulator of developmental disorders, particularly cancer, such a B-cell chronic leukemia.

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18. The composition of any one of claims 13-15 as a marker or modulator of gene expression.

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- 19. The composition of claim 18 as a marker or modulator of the expression of a gene, which is at least partially complementary to said nucleic acid molecule.
- 20. A method of identifying microRNA molecules or precursor molecules thereof comprising ligating 5'- and 3'-adapter molecules to the ends of a size-fractionated RNA population, reverse transcribing said adapter-containing RNA population and characterizing the reverse transcription products.

Fig. 1 A

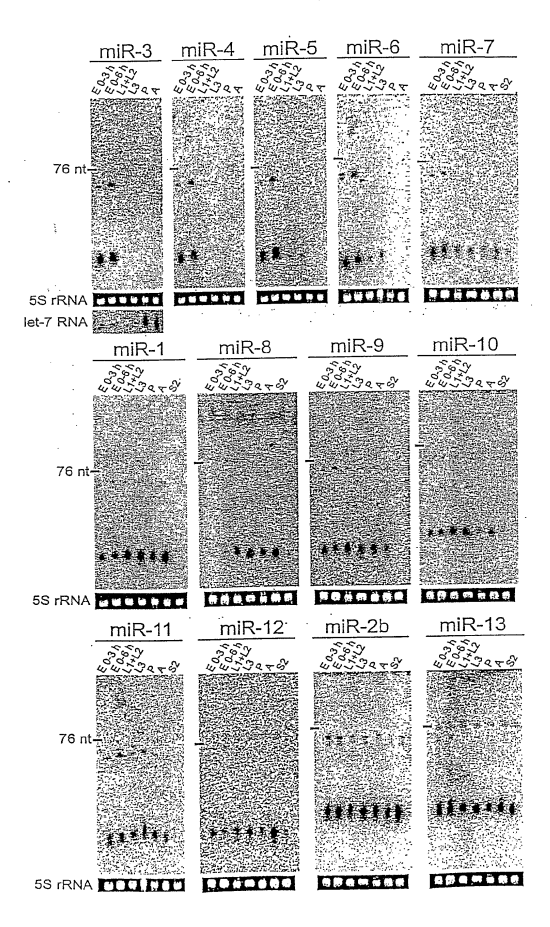


Fig./ B

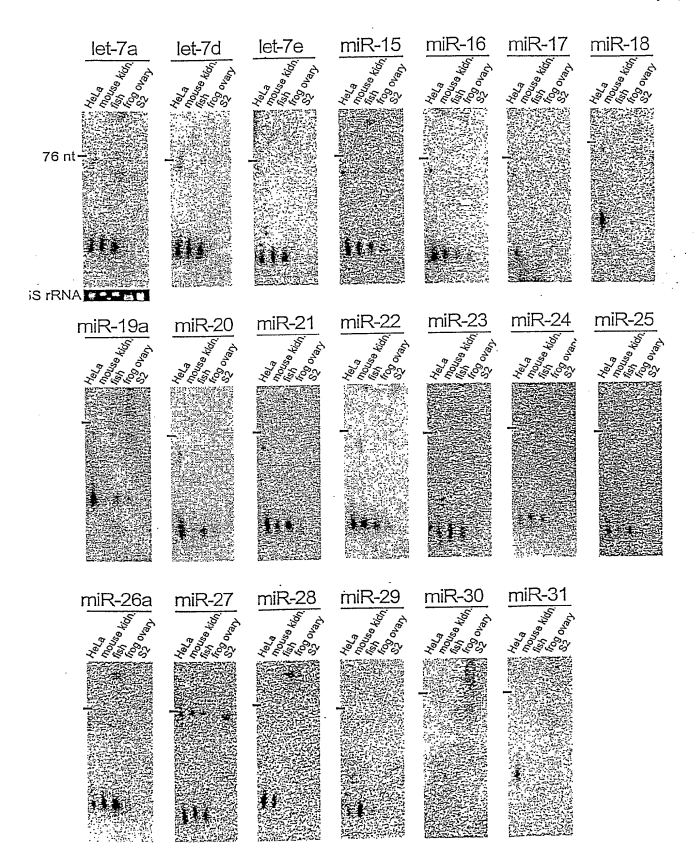


Fig. 2

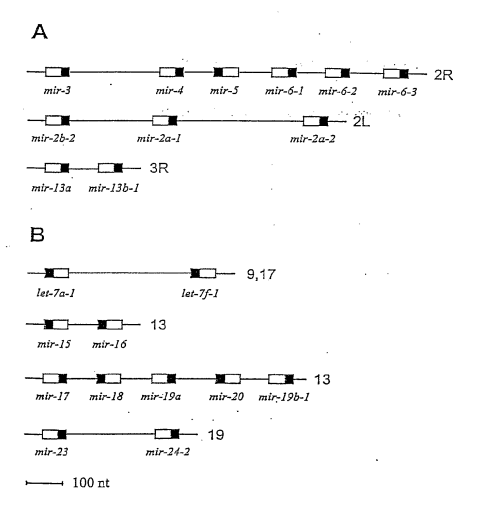


Fig. 3

mir-1	- newtys $\overline{y}$ $\overline{g}$ $y$ yer cores generating $\overline{y}$ $\overline{g}$ $y$ yer 2, and core and $\overline{y}$ $\overline{g}$ $y$ $y$ $\overline{g}$ $y$ $\overline{g}$ $y$	mir-7 5- exerces a central entraces accord according to carry according a contract of carry according a contract of carry according to carry according to carry according to carry according to according to the carry according to t
mir-2a-1	CONTROL RENTAL RECORDERS TO THE COS TO THE C	will-8 additions $\frac{7}{100000}$ $\frac{7}{100}$ $\frac{7}{100000}$ $\frac{7}{100000}$ $\frac{7}{100000}$ $\frac{7}{1000000}$ $\frac{7}{1000000}$ $\frac{7}{1000000}$
mir-2a-2	7 - '73 CCTYC.  2. YDCG YOC GCYGCYGA ACCOYCYGAYG C  7 C GYGYGAGA ACCOYCYGAYG C  7 C GYGYGAGAGAYGYGYGYGYGYGYGYGYGYGYGYGYGYGY	Mit-8 2, ocay nama ayaqcay a cyacay yayaca cy y a yaya ayaya cy y caya yayaca cy y a cyacay yayaca cy y a cyacay yayaca cy y a yaya ayaya cy y a yaya ayaya ayaya ayaya ay yaya ayaya a ayaya ayaya ayayaya ayaya ayayaya ayayaya ayaya ayayaya ayayay
<i>mir-2b-1</i> chr. 2L	C CO	### A PAGE OF THE PROPERTY OF
<i>mir-2b-2</i> chr. 2L clust	EL 7 P DEPARE DATE  2. DECORAGE DECORPORAÇÃO DE CO D  Y - Y DECORAGE DE CODE  Y - Y DECORAGE DATE  Y - COD	mir-1.1 2. Georgian - Georgian
mir-3	$\frac{y}{2}$ $\frac{y}{2}$ $\frac{y}{2}$ $\frac{z}{2}$ COCO $\frac{y}{2}$ COCO $\frac{y}$	mir-12  5' taccord techna town town town can a located to the control town town town town town town town town
mir-4	C $\overline{m}$ $\overline{y}$ $\overline{y}$ $\overline{y}$ $\overline{y}$ $\overline{y}$ CC censence another $\overline{y}$	mir-13a 5' taca along collection and call a collection and call a collection and call a collection and call a collection and c
mir-5	COMES - YNGCCA CO TANGCON GANGGRANDGO A 2.0C TANGCON GANGGRANDGO A C TANGCON A C TANGCON GANGGRANDGO A C TANGCON A	mir-13b-1 5° ccx a recordance exercity and c control of
mir-6-1	CG	mir-13b-2 5. The concentrate detects a process of cher. X and a concentrate detect of cher. X and a co
mir-6-2	$\overline{a}$ $\overline{ac}$ $c$ $c$ $r$ ended $\overline{acatematical}$ $acatematica$	mir-14  5' попосода слад оберал делот дел
mir-6-3	### ##################################	•

Fig. 4

<i>let-7a-1</i> chr. 9,17	$\frac{y_0 - c_0}{\Delta} \qquad \frac{y_0 - c_0}{\Delta} \qquad y_0 - c_$	mir-20	y $yy$ - $a$ $aq$ $aq$ $aq$ $aq$ $aq$ $aq$ $aq$
<i>let-7a-2</i> chr. 11	ם- ס כ yq מככ <u>החכ</u> אהם הככסאכאמתבמא באש ס 2, אפס <u>פאס מאס אפתמתפמאמאתם אמ</u> אה פ הה ס ה	mir-21	S. Desicoscando contro de contro con con con con con con con con con co
<i>let-7a-3</i> chr. 22	а дуасалуалустаную дуасалуда д дуасалуда д доссая с с с с с с с с с с с с с с с с с с	mir-22	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
let-7b	ancea anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy and anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy anceancy and anceancy anceancy anceancy and anceancy anceancy anceancy and anceancy anceancy and anceancy anceancy anceancy and anceancy anceancy and anceancy anceancy and anceancy anceancy and anceancy anceancy and anceancy anceancy and anceancy anceancy anceancy anceancy and anceancy an	mir-23	7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7
let-7c	CA A A A A A A A A A A A A A A A A A	<i>mîr-24-1</i> chr. 9	7         7         2         6         сустан           20         60 </td
let-7d	equation access by converge $\gamma$	mir-24-2 :-chr. 19	7 <u>YCA</u> CYCY AQ YCA AG YCA AG CYCY AQ
let-7e	$y$ CD $q$ - Yangay $c$ 2, CC 600 $\overline{q}$ $\overline$	mir-25	
<i>let-7f-1</i> chr. 9,17	7000	mir-26a	у с - уссс сэс сөвөсү алгауда асаасаудассаад а г, еда ссасаа <u>суувауу суувуулувасда</u> а в й й сого
<i>let-7f-2</i> chr. X	eccentrateracy control of the second of the	mir-26b	Ye C - CC CARA CAPANARA ACARA CARA CARA CARA CARANARA ACARA ACARA ACARA ACARA CARA
mir-1 <i>5</i>	$\frac{y_{12}y_{2}y_{2}y_{2}}{\cos y_{1}}$ and $\frac{y_{12}y_{2}y_{2}y_{2}}{\cos y_{1}}$ and $\frac{y_{12}y_{2}y_{2}y_{2}y_{2}y_{2}}{\cos y_{1}}$ and $\frac{y_{12}y_{2}y_{2}y_{2}y_{2}y_{2}}{\cos y_{1}}$ and $\frac{y_{12}y_{2}y_{2}y_{2}y_{2}y_{2}}{\cos y_{1}}$	mir-27	C C C C COMPANDESSAGE CHEMICAL CA Y  2. CAR CA CA CANAGE CA CANAGE CA Y  Y Y Y  A 0 ACCYC
mir-16	77 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	mir-28	c c c c c c c c c c c c c c c c c c c
mir-17	CO Y <u>nd</u> Y 7 - A CANG  2. CACA DIADACCA <u>DAGACCOY</u> CO YCCAC YAC YC A  2. CACA YMYNGAM YYNGAOCDA CY ACCYO MY CA  CY CY- Y G 4 - YMY	mir-29	2cg         _ ADYA           2v vogazanya         Vccycar         Acma         v           2v vogazanyana         Vccycar         Acma         v           2v vogazanyana         c         acma         v
mir-18	DC $A$ $C$ $C$ $A$ $C$ $C$ $A$ $C$ $C$ $A$ $C$ $C$ $A$ $C$ $C$ $A$ $A$ $C$ $A$ $C$ $A$ $C$ $A$ $A$ $A$ $C$ $A$	mir-30	ور مور وموسومهم <u>حصحمموم</u> محم م دهر <del>محمدممهم حصحممهم محم م</del> ومعمدمهمهم محم محم م
mir-19a	$c$ $\Delta$ $\overline{m}$ $\overline{m}$ $\Delta m$ $\gamma m$ $c$	mir-31	7 7 7 AC CON CCARRICAL CCARR DAY TO TOCCARDAGE GAY A 2. CONTURA GOOTY VAN DOCCARDAGE GAR C 7 G C AT TO TOCKARD CONTURN C 7 G C AT TO TOCKARD C CONTURN C C CONTURN C C CONTURN C C C C C C C C C C C C C C C C C C C
<i>mir-19b-1</i> chr. 13	у д подата саста састосатитела се итела саст у г. суста сатаосатусавання от падаст суст / да подада	mir-32	7 M2 d  CAMADYDYNOROGOGO DOYALDAYCONY C CO C  2, COYOYANDOCYCHA YCDYYCDAYCA G cd c  A
<i>mir-19b-2</i> chr. X	7 Z Zcoa d 2, 757200 датегуратованост од падоста оседура у Сарс	mir-33	$\begin{array}{cccccccccccccccccccccccccccccccccccc$

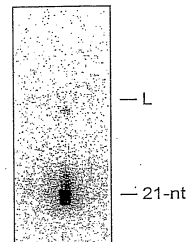
Fig. 5

miR-1a miR-122a

ht kd lv pc sp



ht kd lv pc sp



miR-124a

brain

rbmbcx cb ht ig Iv co si pc sp kd sm st H

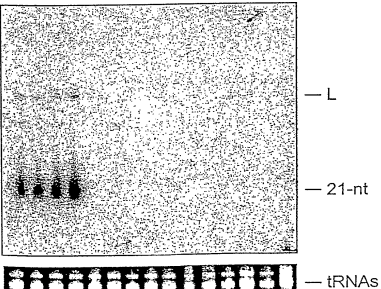


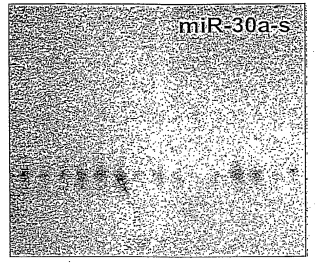
Fig. 5 (cout.)

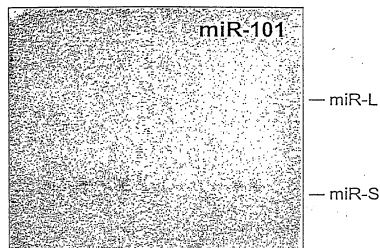
brain

rbmbcxcb ht lg lv co si pc sp kd sm st H



rbmbcx cb ht lg lv co si pc sp kd sm st H







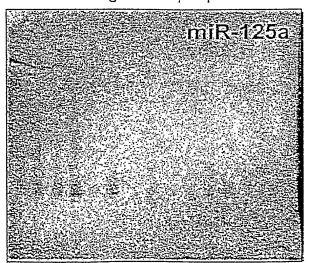
— tRNAs

brain

rbmbcx cb ht lg lv co si pc sp kd sm st H

brain

rbmbcx cb ht lg lv co si pc sp kd sm st H



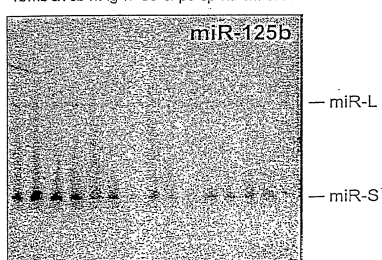


Fig. 5 (cout.)

miR-128

brain

rbmbcxcb ht lg lv co si pc sp kd sm st H

brain

8/46

rbmbcx cb ht lg lv co si pc sp kd sm st H

miR-127

- miR-L

– miR-S

brain

rbmbcxcb ht lg lv co si pc sp kd sm st H

brain

rbmbcx cb ht lg lv co si pc sp kd sm st H

miR-131

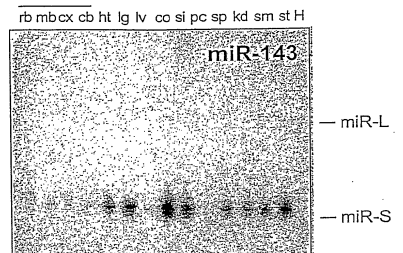
miR-132

— miR-L

- miR-S

Tig. 5 (cout.)

## brain



Tig.6

C. elegans lin-4

D. melanogaster miR-125
M. musculus/H. sapiens miR-125b
M. musculus/H. sapiens miR-125a

UCCCUGAGACCUC--AAG-UGUGA UCCCUGAGACCCU--AACUUGUGA UCCCUGAGACCCU--AACUUGUGA UCCCUGAGACCCUUUAACCUGUGA

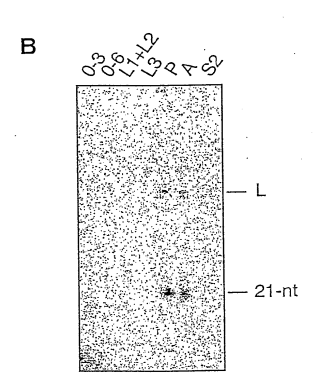


Fig. 7

Sequei AGGUAGUAGGUU AGGUAGUAGGUU AGGUAGUAGGUU	structure	CAC UGGGA GAGGUAGUUGUAUAGUU GUC CCCA C GUG AUCCU UUCUGUCAUCUAACAUAUCAA UAG GGGU A CA A C	GUAVAGUU UAGAAUUAC AA  GUAVAGUU AUC GAGUUGUAUAGUU  UCC UUC AUC UCCGACAUGUCAA  U- G C  AG	GUAUAGUU  GGG GAGGUAGUAGGUUGUAUAGUU  UCC UUCUGUCAUCUAACAUAUCAA  UAGGGUAUC  U	GG U CGGGG GAGGUAGGUUGUGUGGGUU UC GGGCAG \ CGGGG GAGGUAGGUUGUGGGUU UC GGGCAG \ GUCCC UUCCGUCAUCAAAAG CCCGUU A  U AAGGCUC GU	A U <u>U G U</u> GC UCCGGG <u>GAG UAG AGGUUGUAUGGUU</u> GA U C \ GC UCCGGG <u>GAG UAG AGGUUGUAUGGUU</u> GA U C \ CG AGGUUC UUC AUC UCCAACAUGUCAA UU A G C - G GG UC	GGCAUAGU GGAUUCU UUCCGUCGUCCAGC UAUCAA  A UGGAGAACA  UGGAGGAACA  UGGAGGAACA  UGGAGGAACA  UGGAGGAACA  UGGAGGAACA	C CU G U GGA A
	sednence	UGAGGUAGGUUGUAUAGUU	ugagguagguuguaùaguu	UGAGGUAGGUUGUAUAGUU	de ce ce ce co co co	UGAGGUAGGUUGUAUGGUU GC	AGAGGUAGGUUGCAUAGU	C C <u>U</u> <u>G</u> CC GGG <u>GAG UAGGAGGUUGUAUAGU</u> GA

let-7f-1	UGAGGUAGUAGAUUGUAUAGUU	AG <u>U</u> UCAG <u>GAGGUAGAUUGUAUAGUU</u> GU GGGGUAG \ AGUC UUCCGUUAUCAAUA UCCCAUU A CC- GAGGACUUG
let-7f-2	UGAGGUAGUAGAUUGUAUAGUU	U CUGUGGGA <u>GAGGUAGUÁGAUUGUAUUU</u> UUAGGG A GGCACCCU UUCUGUCAUCAAAA GGUUCU C
let-7g	идаддилдилдилсадил	A <u>U</u> A UGAGG A- A A CCC GCC GAGGUAGU GUUUGUACAGUU GUCU UG UACC CCC GCCC UUCCGUCA CGGACAUGUCAA UAGA. AC AUGG CACCCCCCCCCCCCCCCCCCCCCCCCCCCCC
let-7h	UGAGGUAGUAGUGUACAGUU	
let-7i	ugagguagungugggu	U U UGUG CUGGC <u>GAGGUAGUUUGUGC</u> GUU GG CGGGU \ GAUCG UUCCGUCAUCGAACGCG CAA UC GCCCG A U UAGAGGUG - UUAC
miR-1	UGGRAUGURARGARGURUGGAG	A UUUGAGA C A - AUA UUC GCC GUUCCAUGCUUC UUGCAUUC AUA GUU \ GAG CGG C <u>GAGGUAUGAAG AAUGUAAG U</u> AU CGA U
miR-1b	UGGAAUGUAAAGAAGUAUGUAA	A GC AC UGGGA ACAUACUUCUUAUAU CCAUA UGG \ ACUCU <u>UGUAUGAAGAAAUGUA GGU</u> AU AUC C AL449263.5

miR-1c	UGGAAUGUAAAGAAGUAUGUAC	
miR-1d	UGGAAUGUAAAGAAGUAUGUAUU	C GC UGAACC GCUUGGGA ACAUACUUUNUAUAU CCAUA U CGACUUU UGUAUGAAGAAAUGUA GGUAU GAAUC $\frac{\Lambda}{\Lambda}$ CGAAUC
miR-2a-1	UAUCACAGCUUUGAUGAGC	GCUGGGCUC UCAAAG UGGUUGUGA AUGC CGC \ CGAUU <u>CGAG AGUUUC ACCGACACU U</u> ACG GCG U
miR-2a-2	UAUCACAGCCAGCUUUGAUGAGC	AUCU AGC UCAUCAAG UGGUUGUGAUAUG \ UAGG U <u>CG AGUAGUUU ACCGACACUAU</u> AC C A - <u>CG</u> GCAAC
miR-2b-1	UAUCACAGCCAGCUUUGAGGAGC	U UG – A C U CU CAAC UCUUCAAAG UGGC GUGA AUGUUG C GG GUUG $\overline{AGGAGUUUC}$ $\overline{ACCG}$ $\overline{CACU}$ UAUAAC A C $\overline{CG}$ $\overline{G}$ $\overline{A}$ $\overline{AU}$ ACU A
miR-2b-2	UAUCACAGCCAGCUUUGAGGAGC	$\lambda$ - $\lambda$ -
miR-3	UCACUGGGCAAAGUGUGUCUCA	C G U UUCA GAUC UGGGAUGCAU UUGU CAGU AUGU \ CUAG <u>ACUCUGUGUG AACG GUCA U</u> ACA A A <u>G C</u> CUCU

Fig. 7 (court.)

miR-4	auaaagcuagacaaccauuga	U UU C C C GG UU UUGCAAU AGUUUC UGGU GUC AGC UUA UGAUU \ GGUGUUG UUGA <u>AG ACCA CAG UCG AAU A</u> CUGG U C <u>UU A A A A</u> CC
miR-5	аалссаиссиисисаили	UA C RAUCGUUGUGAUAUG \ GC AAAGGAA GAUCGUUGUGAUAUG \ CG UUUCCUU UUAGUGACACUAUAC U CAAUA - AAUCCU
miR-6-1	uaucacaguggcuguucuuuuu	A- UUUA UGUAGAGAAUAGUUGCUGUG UGUA U \ AAAU AUG <u>UUUUUCUUGUCGGUGACAC AU</u> AU A U
miR-6-2	UAUCACAGUGGCUGUUCUUUU	C UU UG C U - G UAACC AAGGGAAC C CUG UGAUAUA UA UU A GUUGG <u>UUUUCUUG</u> <u>G GAC ACUAU</u> AU AU AA A <u>U</u> <u>UC GU</u> - C C A
miR-6-3	uaucacaguggcuguucuuuu	A U AAAC CAAA AGAACGGUUGCUG UGAUGUAG UUG \ GUUU UUUUUUUUUUUUGUCGGUGAC ACUAUAUU AAC U G
miR-7	UGGAAGACUAGUGAUUUUGUUGU	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
miR-8	UAAUACUGUCAGGUAAAGAUGUC	CUGUUC - G C UCCUUU AAGGACAU ACAUCUU ACC GGCAG AUUAGA $\backslash$ UUCCUGUG UGUAGAA UGG CUGUC UAAUCU U CCUG $\mathcal{C}_{-}$ A A A CAAUAU

Fig. 7 (cont.)

		,			T	
GCUA UGUUGGU CUAGCU UAUGA GU A CGAU AUAAU GAAGCCA GAUCGA AUACU CA A U U UUC A GAUA	CU – <u>G U</u> AUACU CCACGU <u>ACC CU UAGA CCGAAUUUGU</u> UUU A GGUGUG UGG GA AUCU GGCUUAAACAGGA G UU A G U AUUUC	U UCU CCC U ACU GCACUUG CAAGAACUU CUGUGA GCG GU U CGUGAGU $\overline{GUUCUUGAG}$ $\overline{GACU}$ $\overline{CGC}$	UG U C - GCCUU UACGGU <u>AGUAU ACAU AGGUACUGGU</u> GU A GUGCCG UCAUA UGUA UUCAUGACCA CA A CA - ACCUA	U C – A UC– CU UACG AACUC UCAAAG GGUUGUGA AUG GA A GUGC U <u>UGAG AGUUUU CCGACACU U</u> AC CU U U <u>U A</u> A UCAU AU	ug- $ug$ - $u$ acu uauu $ug$ - $ug$	UAUU G A GCUA UU AAC CGUCAAAUG CUGUGA UGUGGA U $\overline{\rm UQG}$ GCAGUUUUAC GACACU AUACUU G GU $\overline{\rm A}$
UCUUUGGUUAUCUAGCUGUAUGA	acccuguagauccgaauuugu	CAUCACAGUCUGAGUUCUUGC	UGAGUAUUACAUCAGGUACUGGU	uaucacagccauuuugaugagu	mir-13b-1 uaucacagccauuuugacgagu	UAUCACAGCCAUUUUGACGAGU
mir-9	miR-10	miR-11 (	miR-12	miR-13a	miR-13b-1	miR-13b-2

Fig. 7 (cont.)

			A A	·		
C C GCUU UGUGGAG GAGA GGGGACU ACUGU \ AU <u>AUCCUC CUCU UUUCUGA U</u> GAUA A <u>U U C</u> AAUU	GAGUAAAG <u>UA</u> CCUUG GCAGCACA AUGGUUUGUG UUU \ GGAAC CGUCGUGU UACCGGACGU AAA G AUAAAAAACUC UA	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	AG C - <u>A CG</u> UUA UCUA GCAGCAC GU AAUAUUGG AGAU CAGUUG AUG AGUCGUCGUG CA UUAUGACC UCUA GA A UUAUGACC UCUA	UC C <u>U AGCAGCACG AAUAUUGG G</u> U UGA A GU CACU <u>AGCAGCACG AAUAUUGG G</u> U UGA A CA GUGA UCGUCGUGU UUAUAACC CA AUU U GU UU CA A- AUA	GA Ch- A G G - AUA GUCA AUAAUGU AAGUGCUU CA UGCAG UAG UG \ CAGU UAUUACG <u>UUCACGGA GU ACGUC A</u> UC AC U GG A <u>UG</u> A G - U GUG	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
ucagucuuuuucucuccua	UAGCAGCACAUAAUGGUUUGUG	UAGCAGCACAUCAUGGUUUACA	UAGCACGUAAAUAUUGGCG	only different precursor	ACUGCAGUGAAGGCACUUGU	UAAGGUGCAUCUAGUGCAGAUA
miR-14	miR-15a	miR-15b	miR-16	miR-16	miR-17	mi.R-18

ugn	UGUGCAAAUCUAUGCAAAACUGA	U U GCAG CC CUGUUAGUUUUGCAUAG UUGCAC UACA \ CGUC GG GGU <u>AGUCAAAACGUAUC AACGUG</u> AUGU A C U
ugugo	UGUGCAAAUCCAUGCAAAACUGA	UU
nene	UGUGCAAAUCCAUGCAAAACUGA	CUAC ACAUUG UUACAAUUAGUUUUGCA GG UUUGCAU GCGUAUA A UGUAAU AGUGUUAGUCAAAACGU CC AAACGUG UGUAUAU U
UAAP	UAAAGUGCUUAUAGUGCAGGUAG	C A-GUGCUVAVAGUGCAG UAG UG U GUNG ACU AAGUGCUVAVAGUGCAG UAG UG CGUC UGA UUCACGAGUAUVACGUC AUC AU A A AA - U UG
UAG	UAGCUUAUCAGACUGAUGUUGA	A A D DAA UGUCGGGUAGCUUAUC GACUG UGUUG CUGU G'\ ACAGUCUGUCGGGUAG CUGAC ACAAC GGUA C U
AAG	AAGCUGCCAGUUGAAGAACUGU	U CC - A U CCUG GGC GAG GCAGUAGUUCUUCAG UGGCA GCUUUA GU \ CCG CUC CGU <u>UGUCAAGAAGUU ACCGU CGAA</u> AU CG A U C- ACCC
AUC	aucacauugccagggauuucc	C C $-$ G G CUUC GG CGG UUCCUGG GAUG GAUUUG C C CC ACCUU AGGGACC UUAC CUAAAC U $\overline{\Lambda}$ ACUG

miR-23b	AUCACAUUGCCAGGGAUUACCAC	C U C GUGACU GG UGG GUUCCUGGCA UG UGAUUU U CC ACG <u>ACC UAGGGACCGU AC ACUA</u> AA A <u>C AU</u> AUAGA
miR-24-1	UGGCUCAGUUCAGCAGGAACAG	G G A UA UCUCAU CUCC GU CCU CUGAGCUGA UCAGU \ GAG <u>G CA GGA GACUUGACU GGU</u> CA U A A C CACAUU
miR-24-2	UGGCUCAGUUCAGCAGGAACAG	CC CG CU- AA UU CUCUG UCC UGC ACUGAGCUG ACACAG \ GG <u>GAC AGG ACG UGACUCGGU</u> UGUGUU G <u>A</u> <u>ACU</u> CACA UG
miR-25	cauugcacuugucucggucuga	A AG G UUG UG ACG GGCC GUGUUG AGGC GAGAC G GCAAU CUGG C C CCGG CGUGAC $\overline{\rm UCUG}$ $\overline{\rm C}$ $\overline{\rm CGUUA}$ GGUC U C
miR-26a	UUCAAGUAAUCCAGGAUAGGCU	AGGCC GUG COUCG <u>U CAAGUAA CCAGGAUAGGCU</u> GU G UCCGG CGC GGGCA GUUCAUU GGUUCUAUCCGGUA U G A C -
miR-26b	UUCAAGUAAUUCAGGAUAGGUU	GA - <u>U UC</u> UGUG CCGG CCC AG <u>U CAAGUAAU AGGAUAGGUUG</u> \ GGCC GGG UCG GUUCAUUA UCUUGUCCGAC C AG C - CC
miR-27a	UUCACAGUGGCUAAGUUCCGCU	A A A CUG GG GGCUVAGCUGCU GUGAGCA GG \ GAC CC CG CUUGAAUCGGUGA CACUUGU CU A C C C C

miR-27b	UUCACAGUGGCUAAGUUCUG	AUUG UGAU U AGGUGCAGAGCUUAGCUG GUGAACAG UGG \ UCCAC <u>GUCUUGAAUCGGU CACUU</u> GUU GCC U GA U
miR-28	AAGGAGCUCACAGUCUAUUGAG	C AGACCCUCACAGUCUA UG AGUUA U GGU CUUGCCCUC AGGAGCUCACAGUCUA UG AGUUA U UCA GGACGGGAG UCCUCGAGUGUUAGAU AC UCAGU U C CCUU CU
miR-29a	CUAGCACCAUCUGAAAUCGGUU	uuu c ucaau augacuaaag accacga ucuu a ua <u>uuggcuaaag accacga uc</u> uu a
miR-29b	UAGCACCAUUUGAAAUCAGUGUU	A GU UÜAAÄU AGGA GCUGGUUUCA AUGGUG UUAGAU \\\\\\\\\\\\\\\\\\\\\\\\
miR-29c	UAGCACCAUUUGAAAUCGGuua	
miR-30a-s	miR-30a-s UGUAAACAUCCUCGACUGGAAGC	A GCG C <u>UGUAAACAUCC GACUGGAAGC</u> U GUG A CGU GACGUUUGUAGG CUGACUUUCGG CAC G
miR-30a- as .	cuuucagucgauguuugcagc	A GCG CUGUAAACAUCC GACUGGAAGCU GUG A CGU GACGUUUGUAGG CUGACUUUCGG CAC G  CGU GACGUUUGUAGG CUGACUUUCGG CAC G  C

r.y. 1	_					1
<u>U</u> – ucaua a <u>uguaaacaucc aca cucagc</u> ug c ugcauuuguagg ugu gggucggu a - a ugcgu	UAC <u>U</u> <u>ACA</u> GUGGAA AGA <u>GUAAACA CCU</u> <u>CUCUCAGC</u> U A UCU CAUUUGU GGA GAGGGUCGA G UCU CAUUUGU GGA AAGGAU human	U <u>U</u> GUAAGA GU GUAAACAUC GACUGGAAGCU C CA CG CGUUUGUAG CUGACUUUCGA A U U A	GA <u>G C</u> U- GAA GGAGAG <u>GGCAA AUG UGGCAUAGC</u> <u>G</u> UU C CCUUUC CCGUU UAC ACCGUAUCG CAA U UA A A	U GGAGA <u>UAUUGCACAU</u> <u>ACUAAGUUGC</u> AU G GU A CUUUUAUAGUGUGU UGAUUUAACGUA C CG C	A <u>UU</u> CUGUG <u>CAUUGU</u> <u>G GCAUUG</u> CAUG GG \ GACACUACGUGACA C UGUAACGUAC CC G C UU	A <u>UC</u> <u>U</u> G AAG CAUA <u>ACCCGUAGA CGA CUUGU</u> G UG GUGU UGGGUAUCU GCU GAACGC GC G
UGUAAACAUCCUACACUCAGC	UGUAAACAUCCUACACUCUCAGC	UGUAAACAUCCCGACUGGAAG	GGCAAGAUGCUGGCAUAGCUG	UAUUGCACAUUACUAAGUUGC	GUGCAUUGUAGUUGCAUUG	ACCGUAGAUCCĠAUCUUGU
miR-30b	miR-30c	miR-30d	mir-31	miR-32	miR-33	miR-99a

miR-99b	CACCGUAGAACCGACCUUGCG	C <u>C</u> <u>ACCCGUAGA</u> <u>CGA</u> <u>CU</u> <u>UGCG</u> G GG \ CUGUG UGGGUGUCU GCU GA ACGCC CU C CC GU C ACAC G U
miR-101 .	иасавилсививалива	A GUCCA UCAGUUAUCACAGUGCUG UGCU U AGUCAAUAGUGUCAUGAC AUGG U
miR-122a	UGGAGUGACAAUGGUGUUUGU	GG C UGUCC AGCUG <u>U AGUGUGA AAUGGUGUUG</u> A UCGAUA UCACACU UVACCGCAAAC A UCGAUA A Woodchuck
miR-122b	UGGAGUGUGACAAUGGUGUUUGA	
miR- 122a,b	UGGAGUGUGACAAUGGUGUUUG	
miR-123	CAUUAUUACUUUVGGUACGCG	A A <u>U CG</u> CUG C UGAC GC <u>CAUUAUUACUU UGGUACG</u> UGA A ACUG CG GUAAUAAUGAG GCCAUGC ACU C G C U UCAA- U
miR-124a*	UUAAGGCACGCGGUGAAUGCCA	CUCU G GUGUUCAC GCG CCUUGAUU U GAGA C CGUAAGUG CGC GGAAUUAA C $\frac{A}{A}$ CAUAU

Fig.7 (cont.)

miR-124b	UVAAGGCACGCGGGUGAAUGC	CC A GA UAAUG CUCU GUGUUCAC GCG CCUUGAUU \ GAGA <u>CGUAAGUG CGC</u> GGAAUUAA U AC AC AC
miR-125a	UCCCUGAGACCCUUUAACCUGUG potential lin-4 ortholog	CUGGG <u>U CCUGAGA CCUU ACCUGUGA</u> GG C GGUCCG GGGUUCU GGAC UGGACACU CC G
miR-125b	UCCCUGAGACCCUAACUUGUGA potential lin-4 ortholog	UC CUGAGA CCU ACUUGUGA UAU U CGCAUC GGGUUCU GGA UGAACACU AUG U . CA U C ACA A
miR-126	UCGUACCGUGAGUAAUAAUGC	A U CGCUG C GC CAUUAUUACUU UGGUACG UGA A CG GUAAUAAUGAG GCCAUGC ACU C C
miR-127	UCGGAUCCGUCUGAGCUUGGCU	A U G G C AG CC GCC GCU AAGCUCAGA GG UCUGAU UC \ GG UGG <u>CGG UUCGAGUCU CC AGGCUA</u> AG A C <u>U</u> - <u>G</u> <u>U</u> CU AA
miR-128	UCACAGUGAACCGGUCUCUUU	UUC UAG CU U GUUGGA GGGGCCG CACUGU GAGAGGU U CGACU <u>U CUCUGGC GUGACA CU</u> CUUUA A <u>UUU</u> CAA —— C
miR-129	cunnnnaceencaeecanec	GGAU CUUUUUG GGU GGCUU CU A UCUA GAAAAAC CCA CCCGAA GAC GA A U C UU

Fig. 7 Ccoul.)

Tig. + C	Coul.)					
GA GCUCUUUU ACAUUGUGCU CU \ CU CGGGAAAA UGUAACGUGA GA A GCCAUGU	$egin{array}{lll} \mathbf{G} & \mathbf{C} & \mathbf{G} & \mathbf{U} & \mathbf{A} & \\ \mathbf{GUU} & \mathbf{UUUGGUUAUCUAGCU} & \mathbf{UAUGABG} & \mathbf{GU} & \mathbf{U} \\ \mathbf{CAA} & \mathbf{AA} & \mathbf{UG} & \mathbf{AAGCCAAUAGAUCGA} & \mathbf{AU} & \mathbf{AU} & \mathbf{U} \\ \mathbf{A} & \mathbf{A} & \mathbf{A} & \mathbf{C} & \mathbf{G} \\ \end{array}$	A UUC G- G GGGC ACCGUGGCU GAUUGUUACU UGG \ CCC <u>G UGGUACCGA CUGACAAU</u> GG GCC A . C	A AAUAACCAAAUC U GCUA AGCUGGU AA GG ACCAAAUC U CGA <u>U UCGACCA UU CC UGGUU</u> UAG U	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	UU CUAUGGCUUU AUUCCUAUGUGA \ GGUGCCGAGG UAGGGAUAUACU U U— CGCUCG	C <u>UUU</u> UUCU GAGG <u>ACUC AUUUG UGAUGAUGGA</u> \ CUUCUGAG UAAAC GCUACUU U - UCU CGAA
CAGUGCAAUGUUAAAAGGGC	UAAAGCUAGAUAACCGAAAGU	UAACAGUCUACAGCCAUGGUCGU	uugguccccuucaaccagcugu	UGUGACUGGUUGACCAGAGGGA	UAUGGCUUUUUAUUCCUAUGUGAA	ACUCCAUUUGUUUGAUGAUGGA
miR-130	miR-131	miR-132	miR-133	miR-134	miR-135	miR-136

Fig. 7 (conf.)

G G G COUCGGU AGUA CG \ CUUCGGU ACG GUAUUCGUGG UAAUA CG \ GGAGCU <u>G UGC CAUAAGAAUUCGUU AU</u> UGU GC U	C <u>AGCU GGUGUUGUGAA</u> GGCCG GAG AG C GUUGG CCACAGCACUU 'UCGGC UUC UC A GA UA- CCA	G - <u>U A</u> GUGGC GU UAU <u>UCUA CAG GC CGUGUCU</u> CCAGU \ CA AUGAGGU GUC CG GCGCAGAGGUCG U human U C - GAGGC	CCUG CC <u>GUGGUUUVACCCU UGGUAG</u> G ACG A GGAC GG CACCAAGAUGGGA ACCAUCU UGU U	U U GARG GGG CCAUCUU CCAG GCAGUGUUGG GGUU \ CCC <u>GGUAGAA GGUC</u> <u>UGUCACAA</u> UC UCGA U	AC- A UAA G CCAUAAAGUAG AAGCACUAC CA C GGUAUUUCAUC UUUGUGAUG GU A GUA	AC-CAUAAAGUAG AAGCACUAC CA C GGUAUUCAUC UUUGUGAUG GU A GUA
UAUUGCUUAAGAAUACGCGUAG	AGCUGGUGUUGUGAAUC	ucuacagugucu	AGUGGUUUUACCCUAUGGUAG	AACACUGUCUGGUAAAGAUGG	CAUAAAGUAGAAAGCACUAC	uguaguguuccuacuuuaugg
miR-137	miR-138	miR-139	miR-140	miR-141	miR-142s	miR- 142as*

new	AUAAGACGAGCAAAAAGCUUGU	G C GG C AU UGAC GGCGAGCUUUU GC CG UUAUAC UG \ ACUG UUGUUCGAAAA CG GC AAUAUG AC G G AAUAUG AC G AL049829.4
miR-143	ИСАСАГОСАСИСИАССИСА UUAGAUGAAGCACUGUAG	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
miR-144	UACAGUAUAGAUGAUGUACUAG	G A A A- GU GGCUGG AUAUCAUC UAUACUGUA GUUU G CU <u>GAUC UGUAGUAG AUAUGACAU</u> CAGA A A GU
miR-145	GUCCAGUUUUCCCAGGAAUCCCUU	CUCA GG CAGU UU CCAGGAAUCCCU \ GAGU UC GUCA AA GGUCCUUAGGGG C - UU U A UAGAAU
miR-146	UGAGAACUGAAUUCCAUGGGUUU	C <u>U</u> AGCU <u>GAGAACUGAAUU CAUGGGUU</u> A UCGA UUCUUGACUUAA GUGUCCAG A C-A A
miR-147	GUGUGGAAAUGCUUCUGCC	A- CAA ACA GA AAUCUA AGA CAUUUCUGCACAC CCA \ UUAGAU <u>UCU GUAAAGGUGUGUG</u> GGU C <u>CG UC</u> - ACCGAA AU human
miR-148	UCAGUGCACUACAGAACUUUGU	- A- CC - AGU GAGGCAAAGUUCUG AG CACU GACU CUG \ CUC <u>UGUUUCAAGAC UC GUGA CU</u> GA GAU A AL AC - A AGU human

Fig. 7 (cont.)

tg. 1 (cour.)						
GCUCUG CUC GU UCUUC CUCCC UUU U UCGGGGC GAG CA GGAGG GAGGG GAG C	<u>AC U UG</u> UG CCCUG <u>UCUCCCA CCU GUACCAG</u> CUG \ GGGAUAGGGGGU GGA CAUGGUC GAC C CCA UC	C CA UGUCU CCUG CCUCGAGGAGCU CAGUCUAGUA \ GGAC GGAGUUCCUCGG GUCAGAUCAU A A A-	G A CC CGG C CCGGGCUNCUGU AU CACU GACU GCU U GCCCGGGUUCAAGACA UA GUGA CUGA CGA G	GU A- AAU CAGUG UCAUUUUUGUGAU UGCAGCU GU \ GUUAC <u>AGUGAAACACUG</u> <u>ACGUU</u> GA CG A U <u>AU</u> CC AGU	U - CCU UUU GAAGAUAGGUUA CCGUGU UG UCGC \ UUUUUAUCCAGU GGCACA AC AGUG A U UAAGC UUU	U U A UGGCC CUG <u>UUAAUGCUAAU G G UAGGGG</u> UU \ GACAAUUACGAUUG U C AUCCUCAG U
ucuggcuccgugucuucacucc	UCUCCCAACCCUUGUACCAGUGU	CUAGACUGAGGCUCCUUGAGGU	ucagugcaugacagaacuugg	UUGCAUAGUCACAAAAGUGA	UAGGUUAUCCGUGUUGCCUUCG	UUAAUGCUAAUUGUGAUAGGGG
miR-149	miR-150	miR-151 (	miR-152	miR-153	miR-154	miR-155 [BIC-RNA]

Fig. 7 (cont.)

name	sednence	structure
miR-C1	AACAUUCAACGCUGUCGGUGAGU	U A U CU A GGGAUUCA CCA GG ACA UCAACG GUCGGUG GUUU GGU CC UGU AGUUGC CAGCCAC CAAA U A C AAAACAAA
miR-C2	UUUGGCAAUGGUAGAACUCACA	NO TO TO TO TO TO THE TOTAL TO THE TOTAL TO THE TOTAL TO THE TOTAL TO THE TOTAL TOTA
mir-C3	UAUGGCACUGGUAGAAUUCACUG	G AC GA AC CUGU UAUGGC UGGUA AUUCACUG UGA A GCCAU UAAGUGAC ACU G ACU AA GGAA UG CU
miR-C4	connuaceencaeecaaena	$ \frac{c}{\cos \frac{cu}{66u}} \frac{cu}{66gcuu} \frac{G}{cug}$ $\frac{G}{\cos cu}$
miR-C5	UGGACGGAGAACUGAUAAGGGU	$egin{array}{cccccccccccccccccccccccccccccccccccc$
miR-C6	UGGAGAGAAAGGCAGUUC	AGGAU <u>UGGAG GAAAG CAGUUC</u> CUG GG C UUCCUGGUCUC CUUUC GUCGGGGAC CC C

Fig 7 (cont.)

		<u> </u>		· 1			
structure	ACUUUCCAAAGAAUUC CCUU GGGCUU U UGAAGGGUUUUUUAAG GGAA CCCGAA U	f A $f A$ $f C$ $f CGCUGC$ $f UC$ $f GGCU$ $f CAACACAGGAC$ $f CGGG$ $f CCGA$ $f GUGUGUUCUG$ $f GCUC$ $f C$ $f C$ $f C$ $f C$ $f CCCAGU$	GGGCAUC UVACCGGACAGUG UGGA UC \ CU <u>UGUAG AAUGGUCUGUCAC AU</u> CU AG G C C A C A C A C A C A C A C A C A C A	CA <u>UC</u> UCU <u>CA CCUUGCAUG GGAGGG</u> AGG GU GGGACGUAC CCUCCC C	G G A UCDGAGCUGA UCAGU \ CUCC GU CCU CUGAGCUGA UCAGU \ GAGG CA GGA GACUUGACU GGUCA U	U- <u>UA</u> UU CUGUG <u>GAUAUGUUUGAUAUAU</u> <u>GGU</u> UG \ GACAU UUAUACGAACUAUAUA CUAAU A CC UCAAC UU	
sequence	caaagaauucuccuuuugggcuu	uceueucuueueuuecaecee	UAACACUGUCUGGUAACGAUGU	CAUCCCUUGCAUGGUGGAGGGU	GUGCCUACUGAGCUGACAUCAGU	UGAUAUGUUUGAUAUAUUAGGU	
name	miR-C7	miR-C8	miR-C9	miR-C10	miR-C11	miR-C12	

Fig. 7 (cout.)

Sequence  AACGGAAUCCCAAAAGCAGCU  ACCACAGGGUAGAAUUGACA  ACUGGCCUACAAAGUCCCAGA  ACUGGCCUACAAAGUCCCAGA  ACUGGCCUACAAAGUCCCAGA	structure	AGCGGG AACGGAAUCC AA GCAGCUG GU CU C UCGUCC UUGCUUUAGG UU CGUCGAC UA GA A C	C AGCUCUC ABUUG CAGCCAG G ACUGGAUAC UVAAC GUCGGUC UCCCUC	$egin{array}{cccccccccccccccccccccccccccccccccccc$	A U C A A AGU GAG GCUGGG CUUUG GGGC AG UGAG G CUC U <u>GACCC GAAAC UCCG UC A</u> CUU U C <u>U A G A</u> GUC	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	U AGCAGCACAG AAUAUUGGCA GG G
	sedneuce	CAACGGAAUCCCAAAAGCAGCU AGCGGG AAC	CUGACCUAUGAAUUGACA CUGA	UACCACAGGGUAGAACCACGGA AGGAC GGC AC	AACUGGCCUACAAAGUCCCAG CUC UGACC	UGUAACAGCAACUCCAUGUGGA AUCGGG GUA UAGUCU CAU	U AGCAGCACAGAAAUAUUGGC AGC

Fig 7 C cont.)

structure	GUGAAUU GGU GUUU AUGUUGUUG U CACUUAG CCA CAAA UACAACAAC CAAGUCU	G A CA GA - A GGCUGUGC GGGU GAGGG GUGG GGU AAG G CCGGUA $\overline{CG}$ CCGGUA $\overline{CG}$ CCCGGUA $\overline{CG}$ CCA $\overline{CG}$ CCA UUC C	G - C G UCAUU G UC A AGGGAGA AGG U U CAUU G UC A AGGGAGA AGG U AGUAA U AG U UCUCUUCU UCC G A A A A A A A A A A A A A A A A A A	AAC U C U G G GCC CCAGUGU CAGACUAC UGU CA GAG \ CGG GGUUACA GUCUGAUG ACA GU CUC C AUU C	GGC - C UAGUG GCCGU CAUC UUACUGGGCAG AUUGGA U CGG <u>CA GUAG AAUGGUCCGUC UAAU</u> CU C	U U U U O O O O O O O O O O O O O O O O
ecunes	UAGGUAGUUCAUGUUGUUGG	UVCACCACCUVCVCCACCCAGC	GGUCCAGAGGGGAGAUAGG	CCCAGUGUUCAGACUACCUGUU	UAAUACUGCCUGGUAAUGAUGAC	uacucaguaaggcatuguucu
name	miR-C19	miR-C20	miR-C21	miR-C22	miR-C23	miR-C24

Fig.7 (cont.)

name	sednence	structure .
miR-C25	AGAGGUAUAGCGCAUGGGAAGA	U A- UG C GUUCC UUUUCCUAUGC UAUACUUCUU UGGAU \ CGAGG <u>AGAAGGGUACG AUAUGGAGA</u> A AUCUG U U <u>CG</u> G
miR-C26	UGAAAUGUUUAGGACCACUAG	C U G A C U GGUC AGUGGUUCU GACA UUCA CAGUU UG \ CCA <u>G UCACCAGGA UUGU AAGU</u> GUUAA AC A A U A A C G
miR-C27	UUCCCUUUGUCAUCCUAUGCCUG	U GAGAAUA UGGAC <u>UCCCUUUGUC UCCUA GCCU</u> \ ACUUG AGGGAAACGG AGGGU CGGA U
miR-C28	UCCUUCAUUCCACCGGAGUCUG	CUCUUG CUUCAUUCCAC GGAGUCUG U GAGGAC GAAGUGAGGUG CUUUAGAC G
miR-C29	GUGAAAUGUUUAGGACCACUAGA	U C U G A C U GCC GGUC AGUGGUUCU GACA UUCA CAGUU UG \ CGG CC <u>AG UCACCAGGA UUGU AAGU G</u> UUAA AC A C A <u>A</u> _ C G
miR-C30	UGGAAUGUAAGGAAGUGUGUGG	C U AUAUC CCAGG CCACAUGCUUCUUAUAU C CAUAG \ GGUUU <u>GGUGUGUGAAGGAAUGUA</u> <u>G</u> <u>GU</u> AUC U

Fig 7 (cont.)

				· · · · · · · · · · · · · · · · · · ·
structure	AUC U C G GCC CCAGUGU CAGACUAC UGU UCAG A CGG GGUUACA GUCUGAUG ACA GGUC G A <u>UU</u> _ <u>U</u> GUACAG G	A G CGAAUUUGUG GU C AUAUAU GCCU UAGAA CGAAUUUGUG GU C AUAUA GGGG AUCUU GCUUAGACAC UA C A UGA CA	$\frac{A}{C} = \frac{C}{C} = \frac{A}{C} = C$ $CACA ACC GUAGAU CGA CUUGUG UG UG$ $GUGU UGG UAUCUG GUU GAACAC AC C$ $A A U C - GU$	C U UUG - GGAG AAGG AGGGG CGGGAGGAGC CGGGC G UUCC U <u>CUCC</u> <u>CUCCUCUCG</u> GUUCG C
seguence	UACAGUAGUCUGCACAUUGGUU	CCCUGUAGAACCGAAUUUGUGU a miR-10 variant	AACCCGUAGAUCCGAACUUGUGA A a miR-99a variant	GCUUCUCCUGGCUCUCCUCCUC  AAGG AGGGG GAGGGG  UUCC UCUCC CUCCUC  UCC
пате	miR-C31	mir-C32	mi R-C33	miR-C34

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=	Tio	j.7 (	cout)				33/40					
zebrafish				,								
fugu fish					with slightly diff precursor							
Drosophila				AB003659 diff. Precursor			,					
	apleen				EST A1481799.1 spleen = cerebellum (mammary)			FOUND	found			
	heart						found				•	
	midbrain	toung		,	found	found	found	found	found	:	found	
	cortex	nearly identical precursor	nearly identical precursor			numerous genomic hits	trace#8358704 found 2 nearly ident prec					found in cortex,no db hit
mouse	cerebellum .				nearly ident precursor trace#48311003	num.genomic hits, ident precursor;diff precursor -> EST AI614897	trace#83587042 nearly ident prec		ident precursor genomic DNA	ident. precursor in mmtrace 18713911	genomic hits,no EST	-
	colon	found					found					
	small intes			-								
		num.hits in trace data, 3 families of similar precursors			nearly identical precursor	identical and diff. precursors						
	C.elegans			AF274345 chrX with diff. precursor								
	human	AC007924 chr9 AC087784 chr 17 identical precursor	AP001359 chr11	AL049853 chr22 7	AL049853 chr22	AP001667 chr21	AC007924.3 chr9 AC087784 chrl7 identical	AC018755 chr19	AC007924 chr9 AC007704 chr17	AL592046 chrX	precursor ident. to mouse in AC092045.2 chr3	
	name	let-7a-1	let-7a-2	let-7a-3	let-7b	1et-7c	let-7d	let-7a	10t-7f-1	let-7 <i>f-</i> 2	let-7g	let-7h

Fig. 7	(con	<b>⊢</b> .)			34	l/ <b>46</b>				
J				BF157601.1 with C23 (diff. precursor)						
·	2L, AE003667				21, AE003663	2L, AE003663	2L, AE003620	21, AE003663	2R, AE003795	2R, AE003795
	21.				21,	21,	31,	<u>17.</u>	ж <u>.</u>	2R,
		found	found, but no db hit	trace hits(ntl- 23) trace#91 523974						
		found		found						,
found, supported found by EST BEG61268										
found, su by EST B										
		no mouse hit (only nt1-21)								
9		097405.1 nt i								
precursor ident. to mouse [ALI]383.19]; also ACO48341.22		AL449263.5 chr20 ntl-21		AL449263.5 chr20 ntl-22 (23G)						
let-7i	niR-1	miR-1b	miR-1c	miR-1d	miR-2a-1	miR-2a-2	miR-2b-1	miR-2b-2	mi R-3	miR-4

Fig.	7 Ccon	.F.)				55/46				
						2diff precurs scaffold 3868 and 2417				
2R, AE003795	2R, AE003795	2R, AE00379	2R, AE00379	2R, AE003791	2R, AE003805	3 <b>L, A</b> E003516	AE001574	3R, AE003735	X, AE003499	3R, AE003708
			,				•			
-				uman			·. #0			
				not cloned, but mouse EST predicts precursor similar to human		found	not found, but AC011194 chr.11 predicts diff. precursor			
				precursor s			predicts dif			
·				T predicts		AF155142,1 chr19 diff prec,sligh,diff prec.s in trace	194 chr.11		•	
				ut mouse EE		AF15514 diff prec,s1 prec.s	, but AC011			
				t cloned, E			not found			
				ou						
				ACO03791 chr19 diff.precursor; EST BF37391 again different		AC005316 chr15 AC026701 chr5 each with diff. precursor	AF287967 chrll (HOX B4/B5)			
n : R : S	miR-6-1	miR-6-2	miR-6-3	miR-7	miR-8	mi.R-9	miR-10	miR-11	miR-12	miR-13a

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Tig.7	Ccont	. )			30	/46				
					AL606727 diff precurs					<b>G46757</b> with a U9C
3R, AE003708	X, AE003446	2K, AEM3833					-			
									,	found
			trace#72 137197 prec slig diff	trace#79 105069		found				
			found		found	9			_	
			,			found trace#7910506 9; nearly ident prec. as in human			-	
					genomic hits with 2 slightly diff precur,trace#502					
						found				
						several trace, near ly ident precursor				
-										
			13, AC069475		13, AC069475 interesting leukemia locus	3, NT_005740.6	13, AL138714	13, AL138714	13, AL138714	13, AL138714
miR-13b-1	min-13b-2	miR-14	nir-15a	miR-15b	miR-16	miR-16	miR-17	miR-18	miR-19a	miR-19b-1

73.7	7 Cco	ut.)			<del></del>		<del></del>		r	<del></del> -
								G46757 similar precursor		
			three.	db					Scaffold 4097 different precursor	
		found								
		found	found	tracefol 540691 prec sli diff		found				
			found		-	found	<del>,</del>		found	
					EST AW124037 hypothal,EST AI848465 cerebellum	found.EST AI286629 (thymus): nearly ident. to min.24-1; EST AAI11466 (whole embryo)	procursor	•	AC055818.9, tr found ace#88471973 precursor diff. from human	
·			AKOO8813 (CDNA),prec ident to human					predicted in mouse (EST AIS95464), but not cloned		found,trace#6986 6494,slight.diff precursor
,	found	found		-		found		r AIS95464)		
-	•		AKOO8813 CDNAS, same precursor					mouse (ES)		
			cDNAs from Var. tissues,ide : ntical precursor					predicted in		found
										-
X, AC002407	13, ALI38714	17, AC004686	several highly similar ESTs: AM961681 shown 19, AC020916		XM_072557.1 chr9,also human ESTs,prec nearly ident to mouse	9, AF043896	19, AC020916	7, AC073842 second ident.copy found in chr7	3, AP000497	2, AC021016
miR-19b-2	ni.R-20	1 miR-21	S S A MiR-22 A 1	miR-23a	miR-23b F	9 miR-24-1	miR-24-2	miR-25	miR-26a	miR-26b

Fig.	7 (1	out.)							
				Scaffold 17670.(A- third copy)	Scaffold 17670 has two copies of this RNA			Scaffold 3483,diff precursor	
		<u> </u>		Scaffo 17670. third copy)	Scaf 1767 two copi			Scaf 3483 prec	
								_	
nd						nd		pu	ਪੁਰ
found	<u> </u>				r q	found		found	found
found			trace, EST, nearly ident prec	FOUND	found, supportd by ESTS				found
	maps : 13								
found	found, maps to chr 13 MGSC mmtrace			found	found	found			
found, but no db found, but no found hit mouse			ident or 346733	AC024913.32;d found iff precursor in EST BG342396 (retina)					
found,b db hit mouse			nearly ident precursor trace[2346733 4,EST AC024913.32	ACC24913.32,d iff precursor in EST BG342396 (retina)	found	found		found	
db . m m				3-1-1-8-	¥	<u> </u>	1166.		dp a
put n			8 2 3 4 6				wlth d sor in #85261	723292	but no
found, hit			found, mmtrace#23467334				found with diff precursor in trace #85261735	trace #72329251	found,but no db hit for mouse
			found, AC024913.3 n 2						
found			found AC024 2	found					
						found, ESTs , trace6802 3889 all with 226			
					· <u>- · · · · · · · · · · · · · · · · · ·</u>	foun ,tra 3889 With			
							1		
916	.l tical	32	)4 // shr7 iis iso	AL035109.1 chrl CLUSTER of miR- 29-b and 29-c; miRNA similar to miR-83		ant 23	<i>L</i> 9	srent	All36164.8 chr.6 supported by ESTS (BF594736.1)
19, AC020916	XM_098943.1 chr9 identical precursor	3, AC063932	7, AF017104 second ident.copy CLUSTER, this cluster also consvd in mouse:	AL035209.1 chri CLUSTER of miR- 29-b and 29-c; miRNA similar to miR-83		ly id. Lin 15467.:	6, ALOJS467	human AF159227.6 chr8,different precursor	AL136164.8 chr.6 suppo by ESTs (BF594736.
19,	chr:	n n	7, AFO second ident. found CLUSTE Cluste consvd mouse:	ALO CLUS 29-b MiRN		fold fold s AL035 chr6		human AF159 chr8, precu	AL13 chr. by E (BF5
miR-27 a	miR-27b	miR-28	mir-29a	miR-29b	miR-29c	nearly ident fold in min-30a-s AL035467.23 chr6	mir-30a- as	miR-30b	miR-30c
道	Ë	i ii	n i	ij	i i	별	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	mi	T i a

Fig. 7 (1941)

rg.	7 60	rut.)			T			<del></del> _	<del></del>	
			G44780 with diff.precursor							
Scaffold 3483,diff fold						U53213.1 T.fluviat ilis				Scaffold_ 3295
								-		
found, but no mouse db										
· .										
				trace 4891071		found				
			-	12 ea .	0982	AK021368.1 срия f. eyeball				
					mntrace #92340982	AK021 eyeba				
							abundant but no db hit,except woodchuck X13234			genomic hits (tracef6108 147), no
AF159227.5 chrB	9, AL353732	9, AL35 <u>4797</u>	22, 299716	AP000962.2 chr21,ident to mouse;[similar to miR-10 and miR-51]	AC018755.3 chr.19; [similar to miR- 10 and miR-51]	ALIS8147.17 chr9 diff precursor			·	
miR-30d	miR-31	miR-32	miR-33	miR-99a	mir-99b	miR-101	miR-122a	miR-122b	miR- 122a,b	miR-123

Fig.	7 (con	<u>~+.)</u>				r	r	·		
								with diff fold AC091299.2		
			Scaffold_ 2358	with diff precursSc affold_32 95		Scaffold_ 828,diff_ prec				
slightly diff precursor AC009251 chr2L			found in AC006590.1 1 With diff fold					-		
				found						
found			A22U			found				
most abundant;seve ral trace hits;precurs= cercbellum	found	found	trace#8398570 found with 5			found			found	
most abundant in most cereb.,genomic abundant;seve hits ral trace* (trace*21097008, hits;precurs* 11737241)	found, but no db found hit	genomic hits trace#33921945, 48262259 and more		mmtrace#3521597 and more	hit in trace#79514537	genomic hit troe#51670230	found, but no db hit	mmtrace 68479278	several trace hits,mouse ar155142	hit#86984641
found										
found in 272504.1 chrIV intron,diff										
	ACO21518 chr8,nearly ident chr20 ALO96828.29	ident precur in AC018755.3 chr 19	AP001359.4 chr11 AP001667.1 chr21(chr21 like mouse)		human AL117190.6 chr.14 same precurs as in mouse	ident in ACO16742.10 chr 2;diff prec in ACO16943.7	human AC018662.3 chr7		AC005317.2 chr 15 aligh.diff precursor,but AC026701.6 chr 5 ident	AL137038.5 chr17 prec sligh.diff from mouse
miR-124a*	miR-124b	miR-125a	miR-125b	miR-126	miR-127	miR-128	miR-129	miR-130	miR-131	mir-132

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五:5.7	- L Cou	.+.)								
Scaffold_ 1049;prec u nearly like mouse		Scaffold 2125 with similar precurs		Scaffold_ 18244 nearly ident to mouse/man						
AC093440.1 diff. Precursor									L	
		found						found		found
found			,							
		m va va		4 8 <del>1</del>		0				
found, tracel	trace#6462031	trace#7149523 5,ESTEF780995 .1(kidn.,spleen)(~chr3huma n)	trace#8607175 3	trace#8977454 3,EST (hypothal)A18 52436.1,ident	mouse EST BB528620.2	found, but no mouse hit				
-	<u>च</u> ⊶	្សាល - មុន	1 K	. NO W		H E				
										-
									found	found
							several trace hits; trace#1053	AC002397 chr6	found	several EST AI153235
AL391221.15 chr6 diff. Precursor(ident to rat L33722.1)	AL132709.5 chrl4 similar precursor	AC092045.2 chr3 AC018659.35 chr12 (ident or simil to mouse)	AL117190.6 chr14 ident to mouse	AC027691.1 chrl ,ident to mouse,nearly ident fish	AC006058.1 chr3 precursor diff	AP003065.2 chr11	AC026468.8 chr.16,precurso r nearly ident,	AC006512,12 chr12,precursor slightli diff	AC004687.1 chrl7 BCL3/myc translocation locus,like mouse	
niR-133	miR-134 [	miR-135	min-136	miR-137	miR-138	miR-139	miR-140	miR-141	miR-142s	miR- 142as*

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Fig. 7 (cout.)

found but no db hit	found, but no found found found found	EST AA290206 .1.trace .2143909	found Scaffold EST  BF163346 Similar  1 lung	trace#34 639321	found	found, no db hit	trace#85 955550	10352	3,8845	in colon cumarata ku
	found, b db hit									found in colon, supported.by trace103700445;close match MGSC in chr18 (additional
		found						trace#8472 1065,10352 801	trace#8845 6669	found in cold trace#837094 MGSC in chril
	AC008681.7 chr5	XM 064366.1 precursor nearly ident	AC008681.7 chr5 GGGA;precur nearly like mouse, see 2 positions above	AC008388.7 chr5 diff precursor	AL592549.7	AC010719.4				human chr 17 ACO04477.1, nearly
new	A.miR-143	XI Pu pi miR-144 ne	AC GC GC miR-145 ne pc	A( d):	A) mir-147	miR-148	miR-149	miR-150	miR-151	hu AC miR-152 ne

miR-153	Acoussiz.z chri ident.precursor				found sever. mmtrace 87010874		V	F25.
<u> </u>	AL132709.5 Chr14 nearly miR-154 'identical precursor				found sever. mmtrace 86715639			7 (10
Ţ.	human BIC RNA.AR402776.1 [BIC-RNA] (has U12C)		found; chr 16 mouse					u+.)

Ŧ	٦٠	j.7 (	Con	(. ب			· · · · · · · · · · · · · · · · · · ·							
zebrafish			AL590150.2	AL590150.2										
fugu fish		scaffold_1819	scaffold_967	scaffold_ 967		scaffold_3671				scaffold 2210, diff. precursor			scaffold_2294	
Drosophila						found								
	skin	found							-	,				
	thymus													
	lung	found												
mouse	testes				found								1	
	kidney							A COLUMN TO THE	found, trace	found, trace #78964803	found, trace #61928192	found, cDNA AI286629.1, has C170	found, trace#71 760450	found, trace #88722637
	eye	mouse trace #76647842	mouse trace #88841093	trace #86029980	trace #13885686	trace #87318220	chr16 AC012526.32	trace #86691995						
	spleen													found
S. Constant	Initiali	with different precursors in chr9 AL158075.11,chr1 AL136321.5	chr7 AC084864.2 similar precursor	chr7 ACO84864.2 ident.precursor	similar precurs.in chr7 AC018662.3	chr15 AC069082.9	chr22 AC005664.2 ident.precursor	chrl AL512443.7 similar prec.			chrX AF222696.1 nearly ident. precursor	chrs XM_098943.1 has C17U;prec.nearly identical to mouse		
	паше	miR-C1	miR-C2	miR-C3	miR-C4	miR-C5	miR-C6	miR-C7	miR-C8	mir-c9	miR-C10	miR-C11	miR-C12	miR-C13

Fig. 7 (cont.)

					mouse				Drosophila	fugu fish	zebrafish
name	กนพลก	spleen	eye	kidney	testes	lung	thymus	skin			
miR-C14	chr11 AC000159.6			found, but no db hit						,	3.7
miR-C15	chr16 AC025468.6 nearly ident.precursor			EST BI687377.1, several trace							
miR-C16	chr17 AC003101.1, similar precursor			Found, trace#95 55103							,
miR-C17	chrll AC000159.6, chrl AC103590.2; dlf.prec.			found, trace #87796602						scaffold_152	
m1R-C18				found, trace #47823768 (close to mir- 16)		found		found			
miR-C19	chr17 AC009789.21 cloned from human cell line only			similar precursor in mouse chr11 AC011194.15	or in 11194.15		-			scaffold_ 18334	
miR-C20	chrl AL355310.19 cloned from human cell line only										
miR-C21	chri ACO63952.15 cloned from human cell line only						;			1	
miR-C22	chris AC007229.1; chri AL137157.7 similar precursor; cloned from human cell line only										
miR-C23					777	found			`	scaffold_2210	
miR-C24					trace #69879879						
miR-C25					trace #49754566						ļ
mir-c26	ALI36001 ident. precursor				trace #11977216				·		

Fig. 7 (cont.)

					mouse				Drosophila	fugu fish	zebrafish
name	าบแลก	spleen	eye	kidney	testes	Jung	thymus	skin			
miR-C27	chr9 AL159990.12 identical precursor	ì	trace #91503159							scaffold_ 725	
miR-C28	XM_036612.4, precursor very similar							XM_149012.1		scaffold_ 13664	
miR-C29	chr14 AL136001.6 nearly identical precursor							trace #18453604			
miR-C30	chr6 AL391221.15 similar precursor							trace #84055510		•	
miR-C31	chr9 AC006312.8							trace #89079710 }	-	scaffold_5830	
miR-C32								U77364.1, intronic location Hoxd4 gene		scaffold_82	
miR-C33							· }	trace: #84780544 		scaffold_ 15612	
miR-C34							trace# 72109322			÷	

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